NOV 28 T. 55636 SEARCH REQUEST FORM

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Requester's Full Name: GARY I Art Unit: 1642 Phone I	Nic Kol	F 77581	Data: 11/28/01
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If more than one search is subm	nitted, please prior	itize searches in order of ne	ed. 19189
Please provide a detailed statement of the	search tonic, and descri	be as specifically as possible the sub	ect matter to be searched.
Include the elected species or structures, I utility of the invention. Define any terms	keywords, synonyms, ac s that may have a special	ronyms, and registry numbers, and comeaning. Give examples or relevan	ombine with the concept or
known. Please attach a copy of the cover		and abstract.	
Title of Invention:	1+ tacked		
Inventors (please provide full names):	Attacker	J	
Earliest Priority Filing Date:	119/96		
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For Sequence Searches Only Please inclu appropriate serial number.		(See at	Hacse U
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Date Completed: 12-04-01	Litigation	Lexis/Nexis	, v.,
Searcher Prep & Review Time:	Fulltext	Sequence Systems	

Patent Family

Other (specify)_

Other

32

Clerical Prep Time:

PTO-1590 (8-01)

Online Time: _

above-mentioned detergents was simpler and less time consuming than by the .beta.-naphthol method.

IT 9005-65-6

RL: BIOL (Biological study)
 (in antigen purification)

L11 ANSWER 37 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1962:63281 CAPLUS

DOCUMENT NUMBER: 56:63281
ORIGINAL REFERENCE NO.: 56:12161c-g

TITLE: Chemical nature of mouse histocompatibility

antigens

AUTHOR(S): Davies, D. A. L.

CORPORATE SOURCE: Microbiol. Research Estab., Salisbury, UK

SOURCE: Nature (1962), 193, 34-6

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

Readily dispersible but essentially insol. prepns. of mouse H2 AB histocompatibility antigen (I) can be prepd. from ascitic fluid. was not pptd. by (NH4)2SO4, was irreversibly adsorbed on substituted ionexchanger cellulose, was labile to heat, acid, alkali, ultrasonication, and freeze-drying. An insol. fraction contg. all I formed when C3H mouse ascitic fluid (BP8 tumor) was dialyzed or dild. with more than 1 vol. H2O. All I activity was sedimented in 90 min. at 80,000 g. Sedimentation 4 times from saline and 2 times from H2O removed all serum proteins. Results of differential centrifugation at intermediate speeds suggested gross polydispersity of particle size. The most active fractions were centrifuged to equil. at 100,000 g in a sucrose d. gradient. All activity was in the T4 and T5 sedimentation bands, both giving specific inhibition, but only the T5 band (II) was capable of inducing agglutinating H2 antibody formation when injected into mice of appropriate strains. About 120 mg. of II was obtained from 1000 mice; the compn. was 10% N, 1.0% P, 0.5% S, 60% protein, 3.5% carbohydrate, 35% lipid, 1% hexosamine, and 0.3% sialic acid, and it gave a milky suspension in H2O but flocculated in saline. In the ultracentrifuge II was polydisperse and sedimented rapidly. II was homogeneous by electrophoresis, d.-gradient centrifugation, and immunological techniques. The smallest entity with I activity may still not have been prepd. Materials from different inbred mouse strains may have differences in lipid compn. Prepns. from C3H, C57Bl, and BALB/c mice carry specificities expected from the known distribution of H2 alleles. On purification of antigen the lipid content increased with increasing activity. The protein component had a specificity of its own, other than I, which was not exposed on cell surfaces, whereas I specificity was exposed. I specificity was lost in freeze-drying, treatment with Na dodecyl sulfate, deoxycholate, Tween 20, lysolecithin, and periodate.

(FILE 'MEDITNE, BIOSIS, EMBASE, WPIDS, CONFSCI, SCISEARCH, SICST-EPTUS, JAPIO' ENTERED AT 12:35:06 ON 04 DEC 2001)

38 SEA ABB=ON PLU=ON L6 AND (L10 OR MONOSACCHARIDE OR TRISACCHARIDE OR DISACCHARIDE OR SACCHARIDE)

29 DOP REM 112 (9 DOPPLICATES REMOVED)

L13 ANSWER 1 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD ACCESSION NUMBER: 2001-549774 [61] WPIDS

DOC. NO. CPI:

C2001-163566

TITLE:

Liquid biodegradable block copolymer composition, useful as a drug delivery system for e.g. growth hormones, antibacterial agents, anticancer or

antiinflammatory agents.

DERWENT CLASS:

A96 B05 B07

INVENTOR(S):

CHOI, I; SEO, M

PATENT ASSIGNEE(S):

(SAMY-N) SAMYANG CORP

COUNTRY COUNT:

93

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001045742 A1 20010628 (200161)* EN 37

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU

77 7W

AU 2001025550 A 20010703 (200164)

APPLICATION DETAILS:

	12111 110 1	KIND 		PLICATION	DATE
	2001045742	•		2000-KR1508	20001221
ΑU	2001025550) A	ΑU	2001-25550	20001221

FILING DETAILS:

PATENT NO	KIND	PATENT NO
AU 20010255	50 A Based on	WO 200145742

PRIORITY APPLN. INFO: KR 1999-60349 19991222

AN 2001-549774 [61] WPIDS

AB WO 200145742 A UPAB: 20011024

NOVELTY - A liquid polymeric composition capable of forming a physiologically active substance-containing implant in a living body is new.

DETAILED DESCRIPTION - A liquid polymeric composition capable of forming a physiologically active substance-containing implant in a living body comprises a water-soluble liquid polyethylene glycol derivative, a block copolymer which is insoluble in water but soluble in the polyethylene glycol derivative and an active substance.

INDEPENDENT CLAIMS are included for:

- (1) an implant formed from the composition; and
- (2) processes for preparing the composition.

USE - The composition is useful for forming active substance containing implants for drug delivery when injected into a body. Dwg.0/1

L13 ANSWER 2 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER: 2001-407957 [43] WPIDS

DOC. NO. NON-CPI: N2001-301872

DOC. NO. CPI:

C2001-123491

TITLE:

Oral transmucosal solid dosage form drug delivery

formulation comprises pharmaceutical agent

absorbable into oral mucosal tissue and present in

solid solution with dissolution agent.

DERWENT CLASS:

A96 B05 P32

INVENTOR(S): PATENT ASSIGNEE(S): CROFT, J; ZHANG, H (ANES-N) ANESTA CORP

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001030288 A1 20010503 (200143)* EN

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG

KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ

PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG UZ VN YU

ZA ZW

B1 20010724 (200146) US 6264981

AU 2001010797 A 20010508 (200149)

APPLICATION DETAILS:

PATENT NO KIND	APPLICATION	DATE
WO 2001030288 A1	WO 2000-US28113	20001012
US 6264981 B1	US 1999-428071	19991027
AU 2001010797 A	AU 2001-10797	20001012

FILING DETAILS:

PATENT	NO	KIND		PAT	ENT NO	
AU 200	101079	97 A	Based on	· WO	2001302	88

PRIORITY APPLN. INFO: US 1999-428071 19991027

AN 2001-407957 [43] WPIDS

WO 200130288 A UPAB: 20010801

NOVELTY - An oral transmucosal solid dosage form drug delivery formulation comprises a pharmaceutical agent capable of being absorbed into oral mucosal tissue, and a dissolution agent. The pharmaceutical agent is in solid solution with the dissolution agent. The dissolution agent has a rate in the solvents found in the oral cavity greater than that of the pharmaceutical agent.

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is also included for oral transmucosal delivery of a pharmaceutical agent by providing a drug formulation having a solid pharmaceutical agent in solid solution with a dissolution agent, administering the formulation into a patient's oral cavity, and delivering the pharmaceutical agent by absorption through a patient's mucosal tissue.

USE - The invention is used for oral transmucosal delivery of a pharmaceutically active substance.

ADVANTAGE - The invention produces faster dissolution rates and, accordingly, higher absorption rates of the pharmaceutically

active substance. It can afford better solubility in saliva and mucosal absorption without comprising stability of the solid dosage during storage. Dwg.0/3

L13 ANSWER 3 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER: 2001-244222 [25] WPIDS

DOC. NO. CPI: C2001-073233

TITLE: Pharmaceutical system for improved absorption of

hydrophilic agent includes hydrophilic surfactant and is free of triglycerides.

DERWENT CLASS: A96 B05 B07 D16

INVENTOR(S): CHEN, F; FIKSTAD, D T; PATEL, M V

PATENT ASSIGNEE(S): (LIPO-N) LIPOCINE INC; (CHEN-I) CHEN F; (FIKS-I)

FIKSTAD D T; (PATE-I) PATEL M V

COUNTRY COUNT: 9

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001012155 A1 20010222 (200125)* EN 112

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG UZ VN YU

ZA ZW AU 2000060838 A 20010313 (200134)

US 2001024658 A1 20010927 (200159)

APPLICATION DETAILS:

PATENT NO KIND	Al	PPLICATION	DATE
WO 2001012155 A1 AU 2000060838 A US 2001024658 A1	CIP of US	0 2000-US18807 J 2000-60838 S 1999-375636 S 2000-751968	20000710 20000710 19990817 20001229

FILING DETAILS:

PATENT NO	KIND	PATENT NO
AU 20000608	38 A Based o	n WO 200112155

PRIORITY APPLN. INFO: US 1999-375636 19990817

AN 2001-244222 [25] WPIDS

AB WO 200112155 A UPAB: 20010508

NOVELTY - Pharmaceutical system comprises:

- (1) a dosage form of an absorption enhancing composition comprising at least 2 surfactants, and
 - (2) a hydrophilic therapeutic agent.

The system is free of triglycerides

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is included for the above absorption enhancing composition.

USE - Used for controlling the rate and/or extent of bioabsorption of the therapeutic agent.

In a relative absorption study, a sample preconcentrate solution comprising (in g): 0.30 Cremophor RH40, 0.20 Arlacel 186, 0.18 sodium taurocholate and 0.32 propylene glycol was diluted with standard hypotonic PBS pH 7.4 buffer and spiked with 0.1 mM cold acyclovir, then 0.5 mu l tritiated acyclovir (specific activity 18.9 Ci/mmol). The obtained aqueous isotonic dispersion was perfused through rat intestinal segments and the appearance of the acyclovir in the mesenteric blood was monitored along with disappearance on the luminal side. Results showed that the absorption of acyclovir relative to a plain buffer was 704%.

ADVANTAGE - Bioabsorption of the therapeutic agent is improved Dwg.0/0

L13 ANSWER 4 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER: 2001-168427 [17] WPIDS

DOC. NO. NON-CPI: N2001-121465 DOC. NO. CPI: C2001-050270

TITLE: New lyophilized, rigid,

surfactant free reagent spheres, for use in diagnostic detection of antigens in whole cell

samples.

DERWENT CLASS: B04 D16 S03

INVENTOR(S): GOERTZ, S; HEMMES, P

PATENT ASSIGNEE(S): (SPEC-N) SPECTRAL DIAGNOSTICS INC

COUNTRY COUNT: 93

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001004633 A2 20010118 (200117)* EN 34

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG UZ VN YU ZA ZW

AU 2000060088 A 20010130 (200127)

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 20010046	533 A2	WO 2000-IB967	20000714
AU 20000600)88 A	AU 2000-60088	20000714

FILING DETAILS:

PATENT NO	KIND	PATENT NO
AII 20000600	88 A Based on	WO 200104633

PRIORITY APPLN. INFO: US 2000-616018 20000713; US 1999-353191

19990714

AN 2001-168427 [17] WPIDS AB WO 200104633 A UPAB: 20010328

NOVELTY - Lyophilized, rigid reagent spheres (I),

essentially free of surfactant, comprise a reagent useful

to detect an analyte in a given sample.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

(1) producing (I); and

(2) bioassay using (I).

USE - (I) are particularly used to perform antibody /antigen reactions, e.g. to detect cardiac or bacterial antigens, cytokines, drugs of abuse (or their metabolites), typically for diagnosis of sepsis and myocardial infarction.

ADVANTAGE - Since they are free of lytic surfactants, (I) can be used where the test system includes intact cells, e.g. whole blood or its cell-containing fractions. (I) retain all the advantages of known reagent spheres, i.e. rapid dissolution, ease of handling, adequate mechanical stability and uniform distribution of reagents. Dwa.0/0

L13 ANSWER 5 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD WPIDS

2001-496382 [54] ACCESSION NUMBER:

1997-145374 [13]; 1997-145380 [13]; 2001-496010 CROSS REFERENCE:

[51]

DOC. NO. CPI:

C2001-148984

TITLE:

Reconstituted protein formulation for treating chronic medical condition in mammals comprises

lyophilized mixture of a protein/

antibody and a lyoprotectant.

DERWENT CLASS:

B04 D16

ANDYA, J; CLELAND, J L; HSU, C C; LAM, X M; INVENTOR(S):

OVERCASHIER, D E; SHIRE, S J; WU, S S; YANG, J Y

(GETH) GENENTECH INC PATENT ASSIGNEE(S):

COUNTRY COUNT:

PATENT INFORMATION:

LA PG PATENT NO KIND DATE WEEK US 2001014326 A1 20010816 (200154)* 30

APPLICATION DETAILS:

PATENT NO KIN	D	APPLICATION	DATE
US 2001014326 A	l Cont of Div ex Provisional	US 1995-508014 US 1996-615369 US 1996-29182P US 2001-809511	19950727 19960314 19960729 20010314

PRIORITY APPLN. INFO: US 1996-29182P 19960729; US 1995-508014 19950727; US 1996-615369 19960314; US

2001-809511 20010314

2001-496382 [54] AN WPIDS

1997-145374 [13]; 1997-145380 [13]; 2001-496010 [51] CR

AB US2001014326 A UPAB: 20010924

NOVELTY - An isotonic reconstituted formulation (I) comprising a protein/antibody (at least 50 mg/ml) and a diluent is prepared from a lyophilized mixture of a protein/ antibody and a lyoprotectant. The protein/antibody concentration in the formulation is 2 - 40 times greater than in the

> Searcher : 308-4994 Shears

mixture before lyophilization.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) preparation of (I);
- (2) an article comprising a container holding the lyophilized mixture of the protein and the lyporotectant and instructions for reconstituting the lyophilized mixture with the diluent to the protein concentration of at least 50 mg/ml.

USE - For treating mammals (claimed) having chronic medical conditions.

ADVANTAGE - The formulation is sterile, lyophilized and stable at 30 deg. C for at least 6 months or 1 year. The formulation is stable at 2-8 deg. C for at least 30 days. The formulation retains its physical and chemical stability and integrity on lyophilization and storage. The multi-use formulation facilitates ease of use for the patient, reduces waste by allowing complete use of vial contents and results in significant cost saving for the manufacturer since several doses are packaged in a single vial (lower filling and shipping costs). Dwg.0/19

L13 ANSWER 6 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER: 2001-496010 [54] WPIDS

CROSS REFERENCE: 1997-145374 [13]; 1997-145380 [13]; 2001-496382

[54]

DOC. NO. CPI: C2001-148870

TITLE: Stable isotonic reconstituted formulation useful

for treating allergy, asthma and cancer, comprises

specified amount of antibody and diluent, and is prepared from lyophilized mixture

of antibody and lyoprotectant.

DERWENT CLASS: B04 D16

INVENTOR(S): ANDYA, J; CLELAND, J L; HSU, C C; LAM, X M;

OVERCASHIER, D E; SHIRE, S J; WU, S S; YANG, J Y

PATENT ASSIGNEE(S): (GETH) GENENTECH INC

COUNTRY COUNT: 1

PATENT INFORMATION:

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
US 6267958	B1	US 1996-615369	19960314
	Provisional	US 1996-29182P	19960727

PRIORITY APPLN. INFO: US 1996-29182P 19960727; US 1996-615369 19960314

AN 2001-496010 [54] WPIDS

CR 1997-145374 [13]; 1997-145380 [13]; 2001-496382 [54]

AB US 6267958 B UPAB: 20010924

NOVELTY - A stable isotonic reconstituted formulation (I) comprises 50-400 mg/mL of an **antibody** (Ab) and a diluent, prepared from a **lyophilized** mixture of Ab and a lyoprotectant (L)

which prevents or reduces chemical or physical instability of Ab

upon lyophilization and subsequent storage.

DETAILED DESCRIPTION - A stable isotonic reconstituted formulation (I) comprises 50-400 mg/mL of an antibody (Ab) and a diluent, prepared from a lyophilized mixture of Ab and a lyoprotectant (L) which prevents or reduces chemical or physical instability of Ab upon lyophilization and subsequent storage.

In (I), the molar ratio of (L):Ab is about 100-510 mole (L):1 mole of Ab, and Ab concentration in (I) is 2-40 times greater than Ab concentration in the mixture before lyophilization.

INDEPENDENT CLAIMS are also included for the following:

(1) preparation of (I); and

(2) an article of manufacture (II) comprising a container which holds the lyophilized mixture of an antibody and lyoprotectant which prevents or reduces chemical or physical instability of antibody upon lyophilization and subsequent storage, where the molar ratio of lyoprotectant: antibody is about 100-510 mole lyoprotectant:1 mole of antibody, and instructions for reconstituting the lyophilized mixture with a diluent to an antibody in the reconstituted formulation of about 80-400 mg/ml.

ACTIVITY - Cytostatic; antiallergic; antiasthmatic; antiparasitic.

No supporting biological data is given.

MECHANISM OF ACTION - None given.

No supporting biological data is given.

USE - (I) is useful for treating chronic and acute disorders or diseases such as allergy, parasitic infection, interstitial cystitis, asthma, and cancer of breast, ovary, stomach, endometrium, salivary gland, lung, kidney, colon and/or bladder.

ADVANTAGE - (I) is reconstituted to generate a stable reconstituted formulation having a protein concentration which is significantly higher than the protein concentration in the prelyophilized formulation. (I) is a multi-use formulation and facilitates ease of use for the patient, reduces waste by allowing complete use of viral contents, and results in a significant cost savings for the manufacturer, since several doses are packaged in a single vial (lower filling and shipping costs). Dwg.0/19

DERWENT INFORMATION LTD L13 ANSWER 7 OF 29 WPIDS COPYRIGHT 2001

2001-136863 [14] ACCESSION NUMBER: DOC. NO. CPI:

C2001-040061

TITLE:

Stable aqueous pharmaceutical formulation for treating hemorrhagic shock, thermal injury, stroke,

WPIDS

and myocardial infarction, comprises an

antibody not subjected to prior

lyophilization.

DERWENT CLASS:

B04 D16

LAM, X M; OESWEIN, J Q; ONGPIPATTANAKUL, B; INVENTOR(S):

SHAHROKH, Z; WANG, S X; WEISSBURG, R P; WONG, R L

(GETH) GENENTECH INC PATENT ASSIGNEE(S):

COUNTRY COUNT:

1

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

US 6171586 B1 20010109 (200114)* 56

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
US 6171586	B1 Provisional	US 1997-53087P	19970613 19980612

PRIORITY APPLN. INFO: US 1997-53087P 19970613; US 1998-97171

19980612

AN 2001-136863 [14] WPIDS

AB US 6171586 B UPAB: 20010312

NOVELTY - A stable aqueous pharmaceutical formulation (I), comprising an antibody (Ab) not subjected to prior lyophilization, an acetate buffer of pH 4.8-5.5, a surfactant and a polyol, and lacking sodium chloride, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an article of manufacture (II) comprising a container containing (I); and
- (2) stabilizing Ab in (I), by combining Ab not subjected to prior lyophilization, an acetate buffer of pH 4.8-5.5, a surfactant and a polyol, without sodium chloride.

ACTIVITY - Hemostatic; vasotropic; cerebroprotective; vulnerary; cardiant; antiinflammatory; antiulcer; antirheumatic; antiarthritic; cytostatic.

MECHANISM OF ACTION - Vaccine.

No biological data is given.

USE - (I) is useful for treating hemorrhagic shock, thermal injury, e.g. resulting from burns, stroke including ischemic and hemorrhagic stroke, myocardial infarction, inflammatory disorders such as adult respiratory distress syndrome (ARDS), hypovolemic shock, ulcerative colitis, rheumatoid arthritis and B-cell lymphomas.

Dwg.0/28

L13 ANSWER 8 OF 29 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.DUPLICATE 1

ACCESSION NUMBER:

2001371848 EMBASE

TITLE:

The effects of Tween 20 and sucrose

on the stability of anti-L-selectin during

lyophilization and reconstitution.

AUTHOR:

Jones L.S.; Randolph T.W.; Kohnert U.; Papadimitriou A.; Winter G.; Hagmann M.-L.; Manning M.C.; Carpenter J.F.

CORPORATE SOURCE:

J.F. Carpenter, School of Pharmacy, Univ. of Colorado Hlth. Sci. Center, Denver, CO 80262, United States.

john.carpenter@uchsc.edu

SOURCE:

Journal of Pharmaceutical Sciences, (2001) 90/10

(1466-1477). Refs: 39

ISSN: 0022-3549 CODEN: JPMSAE

COUNTRY:

United States
Journal; Article

DOCUMENT TYPE: FILE SEGMENT:

037 Drug Literature Index

039 Pharmacy

LANGUAGE: English SUMMARY LANGUAGE: English

We have chosen an anti-L-selectin antibody as a model protein to investigate the effects of sucrose and/or Tween 20 on protein stability during lyophilization and reconstitution. Native anti-L-selectin secondary structure is substantially retained during lyophilization in the presence of sucrose (1 or 0.125%). However, aggregation of the protein during reconstitution of lyophilized protein powders prepared without sucrose is not reduced by the presence of sucrose in the reconstitution medium. Aggregate formation upon reconstitution is completely inhibited by freeze drying the protein with sucrose and reconstituting with a 0.1% Tween 20 solution. Tween 20 (0.1%) also partially inhibits loss of native anti-L-selectin secondary structure during lyophilization. However, upon reconstitution the formulations lyophilized with Tween 20 contain the highest levels of aggregates. The presence of Tween in only the reconstitution solution appears to inhibit the transition from dimers to higher order oligomers. Potential mechanism(s) for the Tween 20 effects were investigated. However, no evidence of thermodynamic stabilization of anti-L-selectin conformation (e.g., by Tween 20 binding) could be detected. .COPYRGT. 2001 Wiley-Liss, Inc.

DUPLICATE 2 MEDLINE L13 ANSWER 9 OF 29

2001609128 IN-PROCESS ACCESSION NUMBER:

DOCUMENT NUMBER: 21539437 PubMed ID: 11683251

Effect of moisture on the stability of a TITLE:

lyophilized humanized monoclonal

antibody formulation.

Breen E D; Curley J G; Overcashier D E; Hsu C C; **AUTHOR:**

Shire S J

CORPORATE SOURCE: Pharmaceutical Research and Development, Genentech,

Inc., South San Francisco, CA 94080, USA...

breen.deirdre@gene.com

PHARMACEUTICAL RESEARCH, (2001 Sep) 18 (9) 1345-53. SOURCE:

Journal code: PHS; 8406521. ISSN: 0724-8741.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

IN-PROCESS; NONINDEXED; Priority Journals FILE SEGMENT:

Entered STN: 20011102 ENTRY DATE:

Last Updated on STN: 20011102

PURPOSE: To determine the effect of moisture and the role of the AΒ glass transition temperature (Tg) on the stability of a high

concentration, lyophilized, monoclonal antibody. METHODS: A humanized monoclonal antibody was

lyophilized in a sucrose/histidine/

polysorbate 20 formulation. Residual moistures were from 1 to 8%. Tg values were measured by modulated DSC. Vials were stored

at temperatures from 5 to 50 degrees C for 6 or 12 months.

Aggregation was monitored by size exclusion chromatography and Asp isomerization by hydrophobic interaction chromatography. Changes in secondary structure were monitored by Fourier transform infrared (FTIR). RESULTS: T. values varied from 80 degrees C at 1% moisture to 25 degrees C at 8% moisture, there was no cake collapse and were

no differences in the secondary structure by FTIR. All formulations were stable at 5 degrees C. High moisture cakes had higher aggregation rates than drier samples if stored above their Tg values. Intermediate moisture vials were more stable to aggregation than dry vials. High moisture samples had increased rates of Asp isomerization at elevated temperatures both above and below their Tg values. Chemical and physical degradation pathways followed Arrhenius kinetics during storage in the glassy state. Only Asp isomerization followed the Arrhenius model above the Tg value. Both chemical and physical stability at T > or = Tg were fitted to Williams-Landel-Ferry (WLF) kinetics. The WLF constants were dependent on the nature of the degradation system and were not characteristic of the solid system. CONCLUSION: High moisture levels decreased chemical stability of the formulation regardless of whether the protein was in a glassy or rubbery state. In contrast, physical stability was not compromised, and may even be enhanced, by increasing residual moisture if storage is below the Tg value.

L13 ANSWER 10 OF 29 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER:

2000:366976 BIOSIS

DOCUMENT NUMBER:

PREV200000366976

TITLE:

Lyophilized imaging agent formulation comprising a chemotactic peptide.

AUTHOR(S):

Corbo, Diane C.; Link, Mary Jean M. (1); Williams, N. Adeyink; Tomsho, Michelle L.; Bornstein, Michael;

Solomon, Howard F.; Larsen, Scott K.; Suddith, Robert

CORPORATE SOURCE:

(1) Princeton, NJ USA

ASSIGNEE: Ortho Pharmaceutical Corporation, Raritan, NJ, USA; Johnson-Matthey Inc., West Chester, PA, USA

PATENT INFORMATION: US 6024938 February 15, 2000

SOURCE:

Official Gazette of the United States Patent and Trademark Office Patents, (Feb. 15, 2000) Vol. 1231,

No. 3, pp. No pagination. e-file.

ISSN: 0098-1133.

DOCUMENT TYPE:

Patent English

LANGUAGE: A lyophilized imaging agent formulation and methods for making same are disclosed, such formulations comprise a targeting molecule such as antibody or chemotactic peptide, a linker such as diethylenetriaminepentaacetic acid (DTPA) or succinimidyl 6-hydrazinium nicotinate hydrochloride (SHNH), drying protectant such as mannitol, maltose or tricine, and excipient such as polysorbate 80, in citrate buffer. The formulations of the invention are lyophilized and may be stored stably for extended periods of time. Following reconstitution with diluent, the formulations are administered to a subject for scintigraphic imaging or therapeutic use. Also contemplated is a kit comprising a two-vial system wherein a first vial comprises a lyophilized formulation of imaging agent in the form of a lyophilized cake, and a second vial comprises a pharmaceutically acceptable carrier or diluent.

L13 ANSWER 11 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER:

2000-679727 [66] WPIDS

DOC. NO. CPI:

C2000-206815

TITLE:

Parenteral medicinal compositions containing humanized monoclonal antibody fragments

Searcher : 308-4994 Shears

stabilized by blending with non-ionic

surfactants and sugars at weakly

acidic state, free from restrictions in usage e.g.

storage.

DERWENT CLASS:

B04 D16

INVENTOR(S): PATENT ASSIGNEE(S): KOBAYASHI, M; MORI, A; OKADA, A (YAMA) YAMANOUCHI PHARM CO LTD

COUNTRY COUNT:

92

PATENT INFORMATION:

KIND DATE WEEK LA PG PATENT NO ______

WO 2000066160 A1 20001109 (200066)* JA 20

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK

DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX NO NZ PL PT

RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA

ZW

AU 2000043149 A 20001117 (200111)

APPLICATION DETAILS:

PATENT NO KIND	APPLICATION	DATE
WO 2000066160 A1	WO 2000-JP2784	20000427
AU 2000043149 A	AU 2000-43149	20000427

FILING DETAILS:

PATENT NO	KIND	PATENT NO
AU 20000431	49 A Based on	WO 200066160

PRIORITY APPLN. INFO: JP 1999-121424 19990428

2000-679727 [66] WPIDS AN

WO 200066160 A UPAB: 20001219 AB

NOVELTY - A parenteral medicinal composition containing humanized monoclonal antibody fragments, non-ionic

surfactant and sugars with the pH adjusted to

weakly acidic, is new.

DETAILED DESCRIPTION - A parenteral medicinal composition containing humanized monoclonal antibody fragments, non-ionic surfactants and sugars with pH adjusted to weakly acidic.

INDEPENDENT CLAIMS are also included for the following:

(i) a drug preparation for parenteral application which is the freeze-dried material of the composition; and

(ii) a stabilization method for humanized monoclonal antibody fragments, comprising blending them with non-ionic surfactants and sugars at weakly acidic pH.

ACTIVITY - Anticoagulant.

MECHANISM OF ACTION - Antibody inhibition of platelet aggregation.

USE - The composition is used to inhibit blood platelet

ADVANTAGE - The composition is stable and free from

restrictions in usage e.g. storage, transportation and handling.

DESCRIPTION OF DRAWING(S) - A simplified production process of a humanized monoclonal antibody fragment.

Dwg.1/2

L13 ANSWER 12 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER: 2001-033775 [05] WPIDS

DOC. NO. CPI: C2001-010403

TITLE: New antisense oligonucleotide against dystrophin

pre-mRNA exon 19, useful for treating human Duchenne muscular dystrophy patients with loss of

exon 20 in dystrophin mature mRNA.

DERWENT CLASS: A96 B04 D16 INVENTOR(S): MATSUO, M

PATENT ASSIGNEE(S): (JCRP-N) JCR PHARM CO LTD; (MATS-I) MATSUO M;

(NICH-N) JAPAN CHEM RES CO LTD

COUNTRY COUNT: 26

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

EP 1054058 A1 20001122 (200105)* EN 16

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK

NL PT RO SE SI

JP 2000325085 A 20001128 (200110) 11

APPLICATION DETAILS:

		IND		LICATION	DATE
					20000518
JP	2000325085	A	JΡ	1999-140930	19990521

PRIORITY APPLN. INFO: JP 1999-140930 19990521

AN 2001-033775 [05] WPIDS

AB EP 1054058 A UPAB: 20010124

NOVELTY - Use of an antisense oligonucleotide (I) in the manufacture of a therapeutic pharmaceutical composition (II) or medicament for Duchenne muscular dystrophy (DMD) in a human patient with entire loss of exon 20 in the dystrophin mature mRNA, is new. (I) consists of a 20-50 nucleotide sequence against exon 19 of the dystrophin pre-mRNA.

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is also included for (II) for DMD in patients with entire loss of exon 20 in the production dystrophin mature mRNA.

ACTIVITY - Relaxant.

MECHANISM OF ACTION - Antisense therapy.

Restores reading frame of dystrophin mature mRNA. A specimen of muscular tissue was taken from a patient who lacked exon 20 in dystrophin gene, minced and trypsinized to give isolated cells. The cells were washed and then cultured in a growth medium and then subcultivated. When the proportion of myoblasts reached about 80%, the medium was replaced with Fusion medium to induce differentiation into muscular cells. On the fourth day of induction of differentiation, antisense oligoDNA (200 pmol) was introduced into the cells using LipofectAMINE and further cultured for 3.7 and 10 days. After respective incubations, the cells were subjected to

immunohistochemical staining using an antibody against the C-terminus of dystrophin. It was found that dystrophin staining turned positive in the cells in which no dystrophin staining had initially been detected. In addition, staining with an antibody against the N-terminal region of dystrophin also gave a similar result to that obtained with the C-terminal staining, thus confirming that the produced dystrophin extended from the N-terminus to the C-terminus. RNA was extracted from which cDNA was synthesized and amplification comprising a region covering dystrophin exons 18-21 was carried out. The amplification product was sequenced. Results showed that the product had an amino acid reading frame which had been turned in-frame by a direct connection of the exon 18 sequence to that of exon 21 since the fourth day of culture.

USE - In the preparation of medicament and therapeutic composition for DMD in human patients with loss of exon 20 in mature mRNA (claimed). (I) is also useful for treating a human patient with DMD.

ADVANTAGE - (I) shifts the amino acids reading frame in DMD mRNA having an entire loss of exon 20, from an abnormal out-of-frame position, to an in-frame position which converts the disease to the less severe Becker muscular dystrophy.

Dwg.0/0

L13 ANSWER 13 OF 29 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

ACCESSION NUMBER:

2000084305 EMBASE

TITLE:

Development of a **lyophilization** formulation that preserves the biological activity of the platelet-inducing cytokine interleukin-11 at low

concentrations.

AUTHOR:

Page C.; Dawson P.; Woollacott D.; Thorpe R.;

Mire-Sluis A.

CORPORATE SOURCE:

C. Page, Division of Immunobiology, NIBSC, Blanche Lane, South Mimms, Potters Bar, Hertfordshire, United

Kingdom

SOURCE:

Journal of Pharmacy and Pharmacology, (2000) 52/1

(19-26). Refs: 20

ISSN: 0022-3573 CODEN: JPPMAB

COUNTRY:
DOCUMENT TYPE:
FILE SEGMENT:

United Kingdom Journal; Article 039 Pharmacy

037 Drug Literature Index

LANGUAGE:

English

SUMMARY LANGUAGE:

English

AB Recombinant human interleukin-11 (rhIL-11) is a licensed biological therapeutic product in at least one country and is used to combat thrombocytopenia during chemotherapeutic regimens, as well as undergoing clinical trials for a range of other disorders. Following attempts to lyophilize IL-11 at low concentrations, it was clear that a significant loss of recoverable biological activity occurred. Investigation of a variety of factors, including the type of container in which the rhIL-11 was lyophilized, revealed that surface adsorption to glass was a major factor resulting in loss of activity of rhIL-11 in solution (> 40% reduction after 3 h at room temperature), in addition to losses of activity post-lyophilization. To overcome this problem, different formulations containing combinations of human serum

albumin (HSA), trehalose and Tween-20 have been investigated. Two formulations were successful in entirely preserving the biological activity of rhIL-11 through lyophilization and subsequent reconstitution (potency estimates of formulated relative to original material being .gtoreq.0.97). Accelerated degradation studies, performed at intervals over a six-month period, demonstrated the stability of freeze-dried rhIL-11 using these formulations (predicted annual reduction in potency after storage at -20.degree.C .ltoreq.1.4%). In conclusion, we have developed a working combination of excipients (0.5% HSA, 0.1% trehalose and 0.02% Tween-20 in potassium phosphate buffer (pH 7.4)) to formulate a stable rhIL-11 freeze-dried product in glass containers, with no loss in potency. These findings should facilitate development of low dose rhIL-11 products and be an indicator of caution to those using this and other material with similar physical properties, without taking appropriate precautions to avoid losses through adsorption.

L13 ANSWER 14 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER:

WPIDS

DOC. NO. CPI:

1999-302473 [25]

TITLE:

C1999-088642

Preparation of micro- and nano- particle delivery

system.

DERWENT CLASS:

A96 B04 B07

INVENTOR(S):

PROKOP, A

PATENT ASSIGNEE(S):

(UYVA-N) UNIV VANDERBILT

COUNTRY COUNT:

22

PATENT INFORMATION:

PAT	TENT	NO	KIND	DATE	WEEK	LA	PG
MO	9919	3031	Δ1	19990422	(199925) *	EN	52

RW: AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE

W: AU CA JP

AU 9897991 A 19990503 (199937)

A1 20000726 (200037) EN EP 1021168

R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 9918934 AU 9897991 EP 1021168	A1 A A1	WO 1998-US21455 AU 1998-97991 EP 1998-952243 WO 1998-US21455	19981009 19981009 19981009 19981009

FILING DETAILS:

PATENT		KIND				ENT NO	
AU 9897	7991		Based		WO	9918934	_
EP 1021	1168	A1	Based	on	WO	9918934	

PRIORITY APPLN. INFO: US 1997-62943P 19971009

AN 1999-302473 [25] WPIDS

AB WO 9918934 A UPAB: 20011203

> 308-4994 Searcher : Shears

NOVELTY - A method of making particles useful in drug delivery comprises contacting polyanionic polymers with cations in a stirred reactor so that the polyanions and the cations react to form particles, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) a multicomponent system to generate microparticles, composed of a structural (gelling) polymer and a polymer providing mechanical strength and permeability control;
- (2) a particle made by the above method comprising of multicomponent core anionic polymers and anionic antigen, where all components of the core are incorporated as an integral part of the complex formed with the receiving bath polycations;
- (3) a composition of matter comprising multicomponent core polymers, where corona polymers also include a charged surface modifier (electrostatic stabilizer) of the same charge as the corona polymers, and in which all corona components are incorporated in one step as an integral part of the complex;
- (4) a nonionic polymeric surface modifier (steric stabilizer) as part of the corona multipolymeric system, where all corona components are integrated into the outer polymer structure (shell);
 - (5) a vaccine comprising the particles of (2);
- (6) a composition of matter comprising core anionic polymers and anionic antigens (or plasmid DNA or antisense RNA oligonucleotide), all components being incorporated into the ionically formed complex;
- (7) a method of processing reactor content to remove unwanted residual reactants comprises sedimenting or centrifuging the reactor contents, collecting microparticles or nanoparticles generated, rinsing the particles in excess water, buffer or cryopreservation solution, separating suspension by sedimentation or centrifugation, repeating rinsing and separation steps and reducing volume of the suspension to about 1/100th of the initial volume;
- (8) a method of chemical stabilization of the washed and isolated particles, by reacting particles with a crosslinking agent, rinsing in excess of water, buffer, or cryopreservation solution, separating the particles by sedimentation or centrifugation, repeating the rinsing and separation as needed, and reducing the volume of the suspension;
- (9) a method of cryoprotecting the washed particles, by suspending the particles in a cryoprotective solution, and lyophilising;
- (10) a method of immunization of animals comprising the step of orally delivering an encapsulated antigen in the particles, where the particles are taken up by M-cells in Peyer's patches of the epithelial lining of the upper intestinal tract resulting in an increase in secretory and systemic **antibodies** in blood;
- (11) a method of adjusting the biodegradability of polymeric mixtures, by contacting an enzyme with a polysaccharide, and degrading the substrate at physiological conditions in vivo; and
- (12) a method of introducing an adjuvant to potentiate an immunogenic effect, by administration of the adjuvant as part of a droplet forming polymeric mixture.
- USE Possible uses of the micro- or nano- particulate product range over the fields of pharmaceuticals, proteins, polymers, and colloids, immunology, and biomedical engineering. They include delivery of drugs generally, antigens and vaccines for immunization of humans and other animals, genes (plasmid DNA), and antisense RNA

and DNA oligonucleotides. Some targeting is possible, e.g., by adding mocoadhesive polymers to provide sticking to certain mucosal areas; this applies particularly to the M-cells in Peyer's patches in the epithelial lining of the small intestine, to increase delivery of large molecules, e.g., antibodies. It is stated that the particle production operation can be carried out as a continuous, in addition to a batch process.

L13 ANSWER 15 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER: 1999-290190 [25] WPIDS

DOC. NO. CPI: C1999-085883

TITLE: Inhibiting aggregate formation in reconstituted

protein lyophilizates using potassium

phosphate buffer.

DERWENT CLASS: B04 D16

INVENTOR(S): HELLERBRAND, K; PAPADIMITRIOU, A; WINTER, G
PATENT ASSIGNEE(S): (BOEF) BOEHRINGER MANNHEIM GMBH; (HOFF) ROCHE

DIAGNOSTICS GMBH

A1 20000801 (200137)

COUNTRY COUNT: 3

PATENT INFORMATION:

PAT	TENT	NO		KIND	DATE		W)	EEK		LP		PG	}							
EP	9178	 379		A2	1999	0526	5 (:	1999	925)	* GE	;	11	-							
	R:	AL	ΑT	BE	CH CY	DE	DK	ES	FI	FR G	В	SR	ΙE	ΙT	LI	LT	LU	LV	MC	MK
		NL	PT	RO	SE SI															
ΑU	989	4060)	Α	1999	0610) (1999	934)	1										
ZA	9810	0650)	Α	1999	0728	3 (:	1999	935)	•		21								
CN	1220	0270)	Α	1999	0623	3 (1999	943)	+										
CA	225	4145	5	A 1	. 1999	0522	2 (1999	945)	EN	İ									
JP	112	4089	95	Α	1999	0907	7 (1999	947))		11								
ΑU	7142	264		В	1999	1223	3 (:	2000	011)	•										
BR	980	5021	L	Α	2000	0323	L (:	2000)28)	•										
KR	990	4546	50	Α	1999	0625	5 (:	2000	036)	+										
JP	310	5494	1	B2	2000	1030) (:	2000)57)	+		10								
US	6238	3664	1	B1	2001	0529) (2001	L32)	+										

APPLICATION DETAILS:

MX 9809774

PAT	TENT NO	KIND	APPLICATION	DATE
EP	917879	A2	EP 1998-121684	19981113
ΑU	9894060	A	AU 1998-94060	19981120
ZA	9810650	Α	ZA 1998-10650	19981120
CN	1220270	Α	CN 1998-122531	19981120
CA	2254145	A1	CA 1998-2254145	19981120
JΡ	11240895	Α	JP 1998-332681	19981124
ΑU	714264	В	AU 1998-94060	19981120
BR	9805021	Α	BR 1998-5021	19981123
KR	99045460	Α	KR 1998-49921	19981120
JΡ	3105494	В2	JP 1998-332681	19981124
US	6238664	В1	US 1998-196090	19981119
MX	9809774	A1	MX 1998-9774	19981123

FILING DETAILS:

PATENT NO KIND PATENT NO

AU 714264 B Previous Publ. AU 9894060 JP 3105494 B2 Previous Publ. JP 11240895

PRIORITY APPLN. INFO: EP 1998-102846 19980219; EP 1997-120528

19971122

AN 1999-290190 [25] WPIDS

AB EP 917879 A UPAB: 20011203

NOVELTY - Improved method for inhibiting formation of protein aggregates in a reconstituted **lyophilizate** of a pharmaceutical protein-containing composition comprises dissolving the protein in an aqueous solution that contains potassium phosphate (I) as buffering agent and has potassium to sodium ion ratio 10:1 or greater.

DETAILED DESCRIPTION - The solution is frozen, thawed, divided into containers, each containing a dose for injection, then lyophilized.

INDEPENDENT CLAIMS are also included for the following:

- (1) thawable, solid storage form of a protein, with low content of aggregates, that is essentially amorphous, contains a frozen solution of protein in (I) buffer and has K:Na ion ratio at least 10:1; and
- (2) a pharmaceutical composition comprising a protein in aqueous buffer of pH 6-8 that contains (I) as buffering agent, has K:Na ion ratio at least 10:1 and has buffer concentration 10-300 mM. ACTIVITY - None given.

MECHANISM OF ACTION - None given.

USE - The method is preferably used with **antibodies**, particularly those produced by in vitro cell cultures, but may also be applied to immunotoxins, enzymes, protein hormones etc.

ADVANTAGE - The specified buffer stabilizes proteins during freezing, lyophilization, and low temperature storage, so that when reconstitutued the solution formed is practically free from aggregates and particles. It allows proteins that tend to dimerize or multimerize at neutral pH to be formulated as stable frozen compositions.

L13 ANSWER 16 OF 29 MEDLINE DUPLICATE 3

ACCESSION NUMBER: 1999417802 MEDLINE

DOCUMENT NUMBER: 99417802 PubMed ID: 10486434

TITLE: Niosomes as a novel peroral vaccine delivery system.

AUTHOR: Rentel C O; Bouwstra J A; Naisbett B; Junginger H E

CORPORATE SOURCE: Leiden/Amsterdam Center for Drug Research, Division of Pharmaceutical Technology, P.O. Box 9502, NL-2300

RA, Leiden, The Netherlands.

SOURCE: INTERNATIONAL JOURNAL OF PHARMACEUTICS, (1999 Sep 20)

186 (2) 161-7.

Journal code: DA4; 7804127. ISSN: 0378-5173.

PUB. COUNTRY: Netherlands

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199912

ENTRY DATE: Entered STN: 20000113

Last Updated on STN: 20000113 Entered Medline: 19991215

AB The feasibility to develop a peroral vaccine delivery system based on non-ionic surfactant vesicles (niosomes) was evaluated

using BALB/c mice. Ovalbumin was encapsulated in various lyophilized niosome preparations consisting of sucrose esters, cholesterol and dicetyl phosphate. Two different formulations were compared in this study. The specific antibody titres within serum, saliva and intestinal washings were monitored by ELISA on days 7, 14, 21 and 28 after intragastric administration. Only encapsulation of ovalbumin into Wasag7 (70% stearate sucrose ester, 30% palmitate sucrose ester (40% mono-, 60% di/tri-ester)) niosomes resulted in a significant increase in antibody titres. Administration of ovalbumin and empty niosomes did not exert a similar effect, neither did administration of any control formulation. In contrast to ovalbumin loaded Wasag7 niosomes, application of the more hydrophilic Wasag15 (30% stearate sucrose ester, 70% palmitate sucrose ester (70% mono-, 30% di/tri-ester)) niosome preparations did not result in an increase in antibody titres.

L13 ANSWER 17 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER: 1998-

1998-387644 [33] WPIDS

DOC. NO. NON-CPI:

N1998-302308

DOC. NO. CPI:

C1998-117205

TITLE:

New water soluble vinyl membrane-polymer protein

complexes - useful for producing reagent and

diagnosis kits, e.g. for immunoassays.

DERWENT CLASS:

A14 A96 B04 D16 J04 S03

INVENTOR(S):

AUDEBERT, R; POPOT, J; TRIBET, C

PATENT ASSIGNEE(S):

(CNRS) CENT NAT RECH SCI; (CNRS) CNRS CENT NAT RECH

SCI

COUNTRY COUNT:

19

PATENT INFORMATION:

PATENT	 	DATE		 PG
WO 982			(199833) *	

RW: AT BE CH DE DK ES FI FR GB GR IE IT LU MC NL PT SE

W: JP US

EP 946875

A1 19991006 (199946) FR

R: DE FR GB IT

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 9827434 EP 946875	A1 A1	WO 1996-FR2009 EP 1996-942413 WO 1996-FR2009	19961216 19961216 19961216

FILING DETAILS:

PATENT NO	KIND	PATENT NO
FP 946875	Al Based on	WO 9827434

PRIORITY APPLN. INFO: WO 1996-FR2009 19961216

AN 1998-387644 [33] WPIDS

AB WO 9827434 A UPAB: 19980826

New water soluble vinyl membrane-polymer protein amphiphilic

complexes (A) are claimed, in which the vinyl polymer is of formula (I). R1 = COOM, COOR7, N-pyrrolidonyl, phenyl sulphonate or CONR8R9; M = alkali cation; R7 = sugar residue, polyoxyalkylene (preferably polyoxyethylene) with 4-10 alkylene oxide units or (CH2)t-NR10R11; R8, R9 = H, sugar residue, zwitterionic radical or polyoxyalkylene (preferably polyoxyethylene) with 4--10alkylene oxide units; t = 1-5; R10, R11 = H or 1-4C alkyl; R4-R6 = H or methyl; R2 = COOR12 or CONR13R14; R12 = 6-12C alkyl or alkenyl; R13, R14 = 6-12C alkyl or alkenyl, or one of them may also be H; R3 = COOR15 or CONR16R17; R15 = 1-5C alkyl; R16, R17 = 1-5C alkyl, or one of them may also be H; x,y,z are the percentages of the respective repeating units; x = 20-90%; y = 10-80%; z = 0-60%; the average molecular weight is 500-100,000, preferably 50,000 or less.

USE - Reagent kits and immunological diagnostic kits containing (A) are claimed. (A) can be used: to simplify manipulation of protein solutions; to enable proteins to be studied by methods impossible in the presence of surfactants or lipid membranes (e.g. NMR in liquid media, crystallography or electron microscopy of certain types); to obtain concentrated protein solutions (e.g. 10g/l); for storage in lyophilised form, followed by resuspension by simple addition of water or buffer; in diagnostic systems utilising the membrane proteins as receptors or antigens, e.g. for research into circulating antibodies, both soluble or carried by lymphocytes; and in immunisation and the production of antibodies, and to amplify the immune response. The membrane proteins include enzymes, the complexation of which by (I) has potential industrial use. The absence of surfactant enables many pharmacological test to be simplified, e.g. measurement of the affinity of cellular receptors for drugs.

ADVANTAGE - (A) can be lyophilised or made into concentrated solutions; (A) in lyophilised form and aqueous solutions containing (A) at > 5g/l (preferably 10-500 g/l) are claimed. (A) do not contain surfactants. They are stable for several weeks in solution, degradation being comparable with that of proteins in conventional micellar media. The native form of proteins is preserved. Dwg.0/2

L13 ANSWER 18 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD WPIDS 1998-312175 [27]

ACCESSION NUMBER: DOC. NO. CPI:

C1998-096288

TITLE:

Stable lyophilised composition of monoclonal or polyclonal antibodies, which also contains sugar or amino sugar, amino acid and surfactant,

for use as therapeutic or diagnostic agent.

DERWENT CLASS:

A96 B04 D16

KALLMEYER, G; KLESSEN, C; WINTER, G; WOOG, H INVENTOR(S): (BOEF) BOEHRINGER MANNHEIM GMBH; (HOFF) ROCHE PATENT ASSIGNEE(S):

DIAGNOSTICS GMBH 80

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO KIND DATE LA PG WEEK A2 19980528 (199827)* GE 39 WO 9822136

RW: AT BE CH DE DK EA ES FI FR GB GH GR IE IT KE LS LU MC MW NL

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09/308223
           OA PT SD SE SZ UG ZW
        W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI
           GB GE GH HU IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD
           MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR
           TT UA UG US UZ VN YU ZW
    EP 852951
                 A1 19980715 (199832) GE
        R: DE
    AU 9854841
                 A 19980610 (199843)
    ZA 9710409
               A 19990728 (199935)
                                           39
    EP 941121
                 A2 19990915 (199942) GE
        R: AT BE CH DE DK ES FI FR GB GR IE IT LI LU NL PT SE
    CN 1244805 A 20000216 (200027)
    BR 9713521
                 A ·20000321 (200028)
    KR 2000053328 A 20000825 (200121)
JP 2001503781 W 20010321 (200122)
                                           36
    MX 9904565 A1 20000701 (200134)
                B 20010705 (200143)
    AU 735411
APPLICATION DETAILS:
                                    APPLICATION
                                                     DATE
    PATENT NO
               KIND
    _____
                                     WO 1997-EP6452
                                                     19971119
    WO 9822136
                 Α2
    EP 852951
                 A1
                                    EP 1996-118489
                                                     19961119
                                    AU 1998-54841
                                                     19971119
    AU 9854841
                Α
                                    ZA 1997-10409
                                                     19971119
    ZA 9710409 A
                                    EP 1997-951238
                                                     19971119
    EP 941121
                 A2
                                    WO 1997-EP6452
                                                     19971119
                                    CN 1997-181416
                                                     19971119
    CN 1244805
                 Α
                                    BR 1997-13521
                                                     19971119
    BR 9713521
                                    WO 1997-EP6452
                                                     19971119
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FILING DETAILS:

KR 2000053328 A

JP 2001503781 W

A1

B

MX 9904565

AU 735411

PATENT NO K	IND	:	PATENT NO
AU 9854841 EP 941121 BR 9713521 KR 2000053328 JP 2001503781 AU 735411	A Based A Based W Based	on on on on on on on ous Publ.	WO 9822136 WO 9822136 WO 9822136 WO 9822136 WO 9822136 AU 9854841 WO 9822136

PRIORITY APPLN. INFO: EP 1996-118489 19961119

AN 1998-312175 [27] WPIDS

AB WO 9822136 A UPAB: 19990424

Stable lyophilised pharmaceutical composition of monoclonal or polyclonal antibodies, contains a sugar or amino sugar, an amino acid and a surfactant.

USE - The composition is used as a therapeutic or diagnostic

Searcher: Shears 308-4994

WO 1997-EP6452

KR 1999-704336

WO 1997-EP6452

MX 1999-4565

AU 1998-54841

JP 1998-523210

19971119

19990515

19971119

19971119

19990517

19971119

agent (claimed). Dwq.0/0

L13 ANSWER 19 OF 29 WPIDS COPYRIGHT 2001

DERWENT INFORMATION LTD

ACCESSION NUMBER:

1998-163671 [15] WPIDS

DOC. NO. CPI:

C1998-052837

TITLE:

Humoral immunity restorer - contains lactic acid

bacterium.

DERWENT CLASS:

B04 D16

PATENT ASSIGNEE(S):

(NICH-N) NICHI NICHI SEIYAKU KK

COUNTRY COUNT:

PATENT INFORMATION:

LA PG PATENT NO KIND DATE WEEK JP 10029946 A 19980203 (199815)*

APPLICATION DETAILS:

KIND ' APPLICATION DATE PATENT NO ._____ JP 1996-205213 19960715 JP 10029946 A

PRIORITY APPLN. INFO: JP 1996-205213 19960715

1998-163671 [15] WPIDS

JP 10029946 A UPAB: 19980410 AB

> Humoral immunity restorer contains lactic acid bacterium or its treated product and has an activity of restoring humoral immunity function reduced by a chemical.

ADVANTAGE - A drug of high side effects can be dosed by using the humoral immunity restorer. In an example, Enterococcus faecalis NF-1011 was cultured in a Rogosa liquid medium containing 10 g trypticase, 5 g yeast extract, 3 g tryptose, 3 g monopotassium phosphate, 3 g dipotassium phosphate, 2 g triammonium citrate, 1 g Tween 80, 20 g glucose, 0.2 g cysteine hydrochloride and 5 ml salt solution in 1000 ml water at 37 deg. C

for 10-16 hours. The culture was centrifuged and the microbe body was washed with distilled water twice and suspended in distilled water and heated at 110 deg. C for 10 minutes to prepare dead microbe suspension and then freeze-dried to prepare a dry dead microbe body. An antibody-producing

cell was prepared. The microbe body was mixed with powder CE-2 to 5 % and the mixture was dosed freely to a female mouse. The antibody-producing cell was detected by plaque method. Dwg.0/1

L13 ANSWER 20 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER: 1997-145374 [13] WPIDS

1997-145380 [13]; 2001-496010 [51]; 2001-496382 CROSS REFERENCE:

[54]

DOC. NO. CPI:

C1997-046386

New stable reconstituted protein formulations -TITLE:

prepd. from lyophilised mixt. of protein

and lyoprotectant to provide high protein concn..

B04 D16 DERWENT CLASS:

ANDYA, J; CLELAND, J L; HSU, C C; LAM, X M; INVENTOR(S):

OVERCASHIER, D E; SHIRE, S J; WU, S S; YANG, J Y

Shears 308-4994 Searcher :

(GETH) GENENTECH INC 72 PATENT ASSIGNEE(S):

COUNTRY COUNT:

PATENT INFORMATION:

PAT	ENT	NO	1	KIND) Di	ATE		W]	EEK			LA	P	3							
WO	9704	1801		A1	1:	9970	0213	3 (:	 199	713) * :	EN	48	3							
	RW:	ΑT	BE	CH	DE	DK	EΑ	ES	FI	FR	GB	GR	ΙE	IT	ΚE	LS	LU	MC	MW	NL	OA
		PT	SD	SE	SZ	UG															
	W:	AL	MΑ	AT	ΑU	ΑZ	BB	BG	BR	BY	CA	CH	CN	CZ	DE	DK	EE	ES	FI	GB	GE
		ΗU	IL	IS	JP	KE	KG	ΚP	KR	ΚZ	LK	LR	LS	LT	LU	$rac{r}{\Lambda}$	MD	MG	MK	MN	MW
		MX	NO	NZ	PL	PT	RO	RU	SD	SE	SG	SI	SK	ТJ	TM	TR	TT	UA	UG	UZ	VN
ΑU	9665	5992	?	Α	1:	9970	226	6 (:	199	725)										
ZA	9606	6075	,	Α	1:	9980	325	5 (199	819)#		5	5							
NO	9800	335	5	Α	1	9980	0326	5 (:	199	823)										
EΡ	8459	997		A1	1:	9980	0610) (199	827) :	EN									
	R:	AL	ΑT	BE	CH	DE	DK	ES	FI	FR	GB	GR	ΙE	ΙT	LI	LT	LU	LV	MC	NL	PT
		SE	SI																		
BR	9609	9743	3	Α	1	999(0302	2 (199	915)										
JP	115	L017	0	W	1	999(907	7 (:	199	947)		6	9							
MX	9800	0684		A 1	. 1:	9980	0401	L (:	200	004)										
NZ	3135	503		Α	20	0000	128	3 (:	200	015) ′										
ΑU	716	785		В	20	0000	3080	€ (:	200	022)										
ΑU	2000	010	06	3 A	20	0000	3080	9 (:	200	022) #										

APPLICATION DETAILS:

PAT	TENT NO I	KIND			. AP	PLICATION	DATE
WO	9704801	A1			WO	1996-US12251	19960723
ΑU	9665992	Α			AU	1996-65992	19960723
ZA	9606075	Α			ZA	1996-6075	19960717
NO	9800335	Α			WO	1996-US12251	19960723
					NO	1998-335	19980126
ΕP	845997	A1			EP	1996-925497	19960723
					WO	1996-US12251	19960723
BR	9609743	Α			BR	1996-9743	19960723
					WO	1996-US12251	19960723
JР	11510170	W			WO	1996-US12251	19960723
					JP	1997-507749	19960723
MX	9800684	A1			MX	1998-684	19980123
NZ	313503	Α			NZ	1996-313503	19960723
					WO	1996-US12251	19960723
ΑU	716785	В			AU	1996-65992	19960723
AU	2000010063	3 A	Div	ex	AU	1996-65992	19960723
			_		AU	2000-10063	20000112

FILING DETAILS:

PATENT NO	KIND	PATENT NO
AU 9665992 EP 845997 BR 9609743 JP 11510170 NZ 313503 AU 716785	A Based on Al Based on A Based on W Based on A Based on B Previous Po	WO 9704801 WO 9704801 WO 9704801 WO 9704801 WO 9704801 ubl. AU 9665992 WO 9704801

AU 2000010063 A Div ex AU 716785 PRIORITY APPLN. INFO: US 1996-615369 19960314; US 1995-508014 19950727; ZA 1996-6075 19960717; AU 2000-10063 20000112 AN 1997-145374 [13] WPIDS 1997-145380 [13]; 2001-496010 [51]; 2001-496382 [54] CR AB WO 9704801 A UPAB: 20010927 (A) A stable isotonic reconstituted formulation comprises a protein in an amt. of at least 50 mg/ml and a diluent, which reconstituted formulation has been prepd. from a lyophilised mixt. of a protein and a lyoprotectant, where the protein concn. in the reconstituted formulation is about 2-40 times greater than the protein concn. in the mixt. before lyophilisation. Also claimed are: (B) a stable reconstituted formulation comprising an antibody in an amt. of at least 50 mg/ml in a diluent, which reconstituted formulation has been prepd. from a lyophilised mixt. of an antibody and a lyoprotectant, where the antibody concn. in the reconstituted formulation is about 2-40 times greater than the antibody concn. in the mixt. before lyophilisation; (C) a method for preparing a formulation comprising: (a) lyophilising a mixt. of a protein and a lyoprotectant; and (b) reconstituting the lyophilised mixt. of step (a) in a diluent such that the reconstituted formulation is isotonic and stable and has a protein concn. of at least about 50 mg/ml; (D) an article of mfr. comprising: (a) a container which holds a lyophilised mixt. of a protein and a lyoprotectant; and (b) instructions for reconstituting the lyophilised mixt. with a diluent to a protein concn. in the reconstituted formulation of at least about 50 mg/ml; (E) a formulation comprising a lyophilised mixt. of a lyoprotectant and an antibody, where the molar ratio of lyoprotectant:antibody is about 100-1500 mole lyoprotectant: 1 mole antibody; (F) a formulation comprising anti-HER2 antibody (5-40 mg/ml), sucrose or trehalose (10-100 mM), a buffer and a surfactant; and (G) a formulation comprising anti-IgE antibody (5-40 mg/ml), sucrose or trehalose (80-300 mM), a buffer and a surfactant. USE - The anti-HER antibody formulations

USE - The anti-HER antibody formulations antibody can be used to treat or prevent cancers. The anti-IgE formulations can be used for the treatment or prophylaxis of e.g. IgE-mediated allergic diseases, parasitic infections, interstitial cystitis and asthma.

ADVANTAGE - The reconstituted formulations can have high protein concns. and are stable at 2-8 deg. C for at least 30 days (pref. at 30 deg. C for at least 1 year). Dwg.0/19

L13 ANSWER 21 OF 29 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: 1996:253251 BIOSIS DOCUMENT NUMBER: PREV199698809380

TITLE: Development of stable lyophilized

monoclonal antibody formulations: Effect of

excipients on stability.

AUTHOR(S): Bam, Narendra; Dal Monte, Paul R.; Duddu, Sarma P. CORPORATE SOURCE: Pharm. Dev., SmithKline Beecham Pharm., King of

Prussia, PA 19406 USA

Abstracts of Papers American Chemical Society, (1996) SOURCE:

Vol. 211, No. 1-2, pp. BIOT 143.

Meeting Info.: 211th American Chemical Society National Meeting New Orleans, Louisiana, USA March

24-28, 1996 ISSN: 0065-7727.

DOCUMENT TYPE: Conference LANGUAGE: English

DUPLICATE 4 L13 ANSWER 22 OF 29 MEDLINE

ACCESSION NUMBER: 94365558 MEDLINE

DOCUMENT NUMBER: 94365558 PubMed ID: 8083647

Removal/neutralization of hepatitis A virus during TITLE:

manufacture of high purity, solvent/detergent

factor VIII concentrate.

Lemon S M; Murphy P C; Smith A; Zou J; Hammon J; AUTHOR:

Robinson S; Horowitz B

Department of Medicine, University of North Carolina, CORPORATE SOURCE:

Chapel Hill 27599-7030.

JOURNAL OF MEDICAL VIROLOGY, (1994 May) 43 (1) 44-9. SOURCE:

Journal code: I9N; 7705876. ISSN: 0146-6615.

United States PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

Priority Journals FILE SEGMENT:

ENTRY MONTH: 199410

ENTRY DATE: Entered STN: 19941021

> Last Updated on STN: 19941021 Entered Medline: 19941013

Recent reports have suggested an increased risk of type A viral AB hepatitis in hemophilic patients treated with high purity factor VIII concentrates prepared using ion exchange chromatography coupled

with solvent/detergent treatment for inactivation of viruses. To determine the capacity for removal or inactivation of hepatitis A virus during the factor VIII manufacturing process, human plasma and various factor VIII production intermediates were spiked with cell culture-propagated virus and subjected to scaled

down conditions mimicking the manufacture of solvent/ detergent factor VIII. The combination of antibody

-mediated neutralization, cryoprecipitation, anion exchange

chromatography, and lyophilization in the absence of sucrose resulted in a minimal reduction of 5.5 to 8.55 log10

in the infectivity of hepatitis A virus.

DERWENT INFORMATION LTD L13 ANSWER 23 OF 29 WPIDS COPYRIGHT 2001

WPIDS ACCESSION NUMBER: 1990-361472 [48]

DOC. NO. NON-CPI: N1990-275796 DOC. NO. CPI: C1990-157107

Lyophilised immuno assay reagent contg. TITLE:

dextrin or trehalose stabiliserell

culture medium - to prevent agglomeration, immuno reactive material and lig. organic auxiliary,

easily dispersed in aq. medium.

DERWENT CLASS: A89 B04 D16 S03

COLE, F X INVENTOR(S):

(HYGE-N) HYGEIA SCI INC; (HYGE-N) HYGEIA SCIENCES PATENT ASSIGNEE(S):

LTD

COUNTRY COUNT: 16

PATENT INFORMATION:

PAT	TENT NO	KIND	DATE	WEEK	LA	PG
WO	9013637	 А	19901	115 (19904	8) *	31
			DE DK	ES FR GB I	T LU NL	SE
	W: CA	-	10000	010 /1000		
EР	470192			212 (19920 FR GB IT L		SE
US				407 (19921		
				218 (19931		31
				311 (19952		_
EΡ				008 (19974		
חב				ES FR GB I 113 (19975		NT 2F
				216 (19981		
				111 (19985		7

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
EP 470192	A	EP 1990-908003	19900416
US 5102788	A	US 1989-344575	19890428
JP 05500854	W	JP 1990-506814	19900416
		WO 1990-US2064	19900416
EP 470192	A4	EP 1990-908003	
EP 470192	B1	EP 1990-908003	19900416
		WO 1990-US2064	19900416
DE 69031561	E	DE 1990-631561	19900416
		EP 1990-908003	19900416
		WO 1990-US2064	19900416
ES 2110415	Т3	EP 1990-908003	19900416
JP 2823353	B2	JP 1990-506814	19900416
		WO 1990-US2064	19900416

FILING DETAILS:

PATENT NO KIND PATENT NO							
JP 05500854	W Based on	WO 9013637					
EP 470192	B1 Based on	WO 9013637					
DE 69031561	E Based on	EP 470192					
	Based on	WO 9013637					
ES 2110415	T3 Based on	EP 470192					
JP 2823353	B2 Previous Publ.	JP 05500854					
	Based on	WO 9013637					

PRIORITY APPLN. INFO: US 1989-344575 19890428

AN 1990-361472 [48] WPIDS

AB WO 9013637 A UPAB: 19930928

Lyophilised mmixt. for immunoassay comprises (1) at least one dispersible immunoreactive component (I); (2) at least one normally liq. organic component (II) which enhances the performance of the assay, both of these homogeneously distributed throughout the mixt., and (3) sufficient sugar, i.e. dextrin or trehalose, to prevent aggregation of (II), thus maintaining homogeneity.

(I) is an antibody conjugate (AbC), esp. an Ab-gold sol. particle; Ab-solid carrier (esp. latex) particles or an Ab-enzyme conjugate. (II) is an agent which enhances binding, pref. a nonionic, water-soluble polymer (IIa) and/or a surfactant (IIb).

USE/ADVANTAGE - Incorporation of (III) improves shelf life (even when stored under hot, humid conditions) and subsequent dispersion of the mixt. in aq. media for carrying out the immunoassay. The mixt. is easily reconstituted to produce a system contg. all ingredients necessary for an assay. The mixts. are used to assay antibodies, antigens, etc. e.g. human chorionic gonadotropin (hCG) or Neisseria gonorrhoea, esp. in sandwich immunoassay for home and clinical diagnostic applications. They allow such tests to be done by unskilled users without special equipment.

0/0

ABEQ US 5102788 A UPAB: 19930928

Lyophilised mixt. of immunoassay reagents comprises a homogeneously-distributed dispersible immunoreactive component(s), a liq. organic component homogeneously distributed in the mixt. to enhance the performance of the immunoassay, and a sugar, dextrin or trehalose, to prevent agglomeration of the organic components during storage and aid their dispersion in aq. medium during assay.

The immunoreactive component may be an antibody conjugate or antibody-enzyme conjugate. The organic component which enhances binding may be a nonionic water-soluble polymer e.g. polyethylene glycol, PVA, polyvinyl pyrrolidone or dextran and may include a water-soluble nonionic surfactant e.g. a polyethylene glycol, p-isooctyl phenyl ether cpd.

ADVANTAGE - A storage-stable immunoassay system of long shelf life is provided.

ABEQ EP 470192 B UPAB: 19971113

A lyophilised mixture for use in an immunoassay procedure comprising: at least one dispersible immunoreactive component distributed homogeneously throughout said mixture; at least one organic component distributed homogeneously throughout said mixture, said organic component normally being a liquid at the conditions under which the mixture is stored or retained prior to use and having a property which enhances the performance of the immunoassay by its presence, and a sugar comprising dextrin or trehalose, said dextrin or trehalose being present in said mixture in sufficient quantity to prevent agglomeration of the organic component and thus maintain the homogeneity of the mixture to thereby facilitate storgage and shelf life of the mixture and the eventual dispersion of the mixture in an aqueous medium for conduct of the immunoassay procedure.

Dwg.0/0

L13 ANSWER 24 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD ACCESSION NUMBER: 1990-201016 [26] WPIDS

ACCESSION NUMBER: DOC. NO. NON-CPI:

N1990-156415

DOC. NO. CPI:

C1990-087053

TITLE:

Lyophilised mixt. for enzyme immunoassay - contg. antibody enzyme conjugate, binding enhancing agent, water-soluble nonionic surfactant and sugar cpd. to give elevated temp. stability.

DERWENT CLASS:

A96 B04 D16 S03

INVENTOR(S):

BAHAR, I; BLOCK, E; CICIA, N J; COLE, F; COSEO, M;

EATON, C A; JONES, W; SIGILLO, E

PATENT ASSIGNEE(S): COUNTRY COUNT: (HYGE-N) HYGEIA SCIENCES LTD

DAMENT THEODIA TO

-

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

US 4931385 A 19900605 (199026)*

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
US 4931385	A	US 1988-275656	19881121

PRIORITY APPLN. INFO: US 1983-473907 19830310; US 1985-747605 19850624; US 1988-275656 19881121

AN 1990-201016 [26] WPIDS

AB US 4931385 A UPAB: 19930928

In a lyophilised mixt. for use in an enzyme immunoassay, the mixt. comprising an antibody-enzyme conjugate and buffer salts, the improvement is that the enzyme in the conjugate comprises peroxidase and the mixt. further comprises: (a) a binding enhancing agent from PEG, PVA, polyvinyl pyrrolidone (PVP) and dextran; (b) a water-soluble nonionic surfactant in amt. sufficient to provide an appropriate amt. of detergency without having a deleterious effect on the conjugate; and (c) dextrin or trahalose in amt. sufficient to prevent discernible concn. gradients of the components in the mixt.; the lyophilised mixt having the property of preserving the antibody reactivity and the immunologic binding specificity of the conjugate even if the mixt. is exposed to temps. of 80-120 deg.F prior to its use in the immunoassay. Diagnostic kit for ELISA for detection of an antigen in a sample, suitable for home diagnostic applicn. under ambient temp. conditions, comprises the following separately contained components: (1) a solid support precoated with a first antibody and subsequently treated with a blocking soln. comprising a mixt. of a blocking agent and a water-soluble sugar, the blocking agent being BSA, gelatin, milk protein or non-specific IgG antibody; (2) a vial of the above lyophilised mixt.; (3) a measuring dispenser for the sample to be assayed; (4) a container comprising a soln. of a buffer and a peroxide; and (5) a container comprising a soln. of a chromogenic substrate of the peroxidase and a solvent; the components being operable at 15 deg.C to less than 37 deg.C, pref. 15-28 deg.C.

USE/ADVANTAGE - Useful for detection of e.g. human chorionic gonadotropin (LCG) in urinee to detect pregnancy, gonococcal bacteria, luteinising hormone, etc. The assay/kit can be used in the home or physician's office and by unskilled personnel. No specialised appts. is needed. Incubation and washing steps are reduced, thus saving time.

0/0

L13 ANSWER 25 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD ACCESSION NUMBER: 1989-339969 [46] WPIDS

DOC. NO. CPI:

C1989-150711

TITLE:

Homogeneous dimeric macrophage column stimulating factor - and storage stable formulations free of high mol.wt. and stable to prolonged storage.

A96 B04 D16

DERWENT CLASS: INVENTOR(S):

AMPHLETT, G; MORIN, S H; MORRIS, J P; SCHRIER, J A; WILLIAMS, D F

PATENT ASSIGNEE(S):

(GEMY) GENETICS INST INC

COUNTRY COUNT:

PATENT INFORMATION:

PAT	TENT	NO	KIND	DATE		WEEK		LA	PG
WO	8910	0407	A	19891	L102	(1989	946)	* EN	
	RW:	AT, BE	CH I	DE FR	GB 3	T LU	NL :	SE	
	W:	AU JP	KR	-					
AU	8935	5384	Α	19891	L124	(1990	16)		
EΡ	4110	033	Α	19910	206	(1991	106)		
	R:	AT BE	CH I	DE FR	GB 3	IT LI	LU 1	NL SE	
JP	0350	04008	W	19910	905	(1991	142)		
KR	9301	1303	B1	19930)225	(1994	117)		
ΕP	4110	033	B1	19940	907	(1994	134)	EN	6
	R:	AT BE	CH I	DE FR	GB I	IT LI	LU 1	NL SE	
DE	6891	L8091	E	19941	L013	(1994	140)		
		76178							5
		0269						#	
US	5888	3495	Α	19990	330	(1999	920)		

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
EP 411033	 А	EP 1989-905303	19890427
JP 03504008	W	JP 1989-505062	19890427
KR 9301303	B1	WO 1989-US1694	19890427
		KR 1989-702462	19891227
EP 411033	B1	EP 1989-905303	19890427
		WO 1989-US1694	19890427
DE 68918091	E	DE 1989-618091	19890427
		EP 1989-905303	19890427
		WO 1989-US1694	19890427
JP 07076178	B2	JP 1989-505062	19890427
		WO 1989-US1694	19890427
CA 1340269	С	CA 1989-601348	19890531
US 5888495	A Cont of	US 1988-197499	19880523
		US 1992-931551	19920818

FILING DETAILS:

PAT	TENT NO	KIND	P	ATENT NO
EP	411033	B1 Based	lon W	8910407
DE	68918091	E Based	l on El	411033
		Based	lon Wo	8910407
JP	07076178	B2 Based	l on J	03504008
		Based	lon Wo	8910407

19880523; US 1988-187802 PRIORITY APPLN. INFO: US 1988-197499

19880429; CA 1989-601348 19890531; US 1992-931551 19920818

AN 1989-339969 [46] WPIDS

AB WO 8910407 A UPAB: 19930923

Homogeneous dimeric M-CSF, characterised by a single peak when assayed by gel filtration chromatography using a gel filtration matrix having a fractionation range of 10-1500 KD, is new. Also claimed is a storage stable lyophilised m-CSF formulation comprising 0.1-2.0% of m-CSF, 0.1-9% of a polyoxyethylenic non-ionic surfactant, 65-75% glycine, 17-21% sucrose, and 4-7% of a buffering agent, having a pH of 6 upon reconstitution.

USE - m-CSF is a regulatory glycoprotein that stimulates hematoporetic cell proliferation and differentiation. When m-CSF is applied to an in vitro colony stimulating assay it results in the formation of predominantly nonocytic lineage type colonies. m-CSF may be used in activating nature white cells in cases of serious infection or as a therapeutic leukopenia. It may also be used for killing tumour cells either alone or by coadministering it with certain antibodies directed to tumour-associated antigens. 0/0

ABEQ EP 411033 B UPAB: 19941013

A storage stable lyophilised M-CSF formulation of a homogeneous dimeric M-CSF, which is characterised by a single peak when assayed by gel filtration chromatography using a gel filtration matrix having a fractionation range of about 10-1500 kD, comprising about 0.1-10% by weight of M-CSF, about 0.5-20% by weight of a pharmaceutically acceptable polyoxyethylenic non-ionic surfactant, about 40-75% by weight glycine, about 15-40% by weight sucrose and to about 25% by weight of a pharmaceutically acceptable buffering agent, having a pH of about 6 upon reconstitution.

Dwg.0/0

L13 ANSWER 26 OF 29 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: 1989:160335 BIOSIS

DOCUMENT NUMBER: BA87:82436

TITLE: DETECTION OF CMV ANTIBODY USING

FREEZE-DRIED ERYTHROCYTES BY AN INDIRECT HEMAGGLUTINATION TEST.

AUTHOR(S): WANG M; ZHANG Y; GONG Y

CORPORATE SOURCE: DEP. MICROBIOL., ANHUI MED. UNIV., HEFEI.

SOURCE: CHIN J VIROL, (1988) 4 (3), 239-243.

CODEN: BIXUEA.

FILE SEGMENT: BA; OLD LANGUAGE: Chinese

AB This paper reported a study on the methodology for the detection of cytomegalovirus antibody using freeze-

dried CMV antigen-sensitized sheep RBC (SRBC) by an indirect

hemagglutination test. Freeze-drying of CMV

antiqen-sensitized SRBC was carried out in PBS (pH 7.0) containing

5% heat-inactivated normal rabbit serum and 10% sucrose.

The main obstacle to the use of such SRBC was a nonspecific

agglutination occurring after lyophilization of

CMV-sensitized SRBC. This obstacle can be overcome by adding Tween 20 (1/1,000,000) to the phosphate-buffered saline in which the CMV-sensitized SRBC were resuspended for use. The method we established has fairly good reproducibility. The specificity of

the method was the same as ELISA and the sensitivity was higher than

the CF.

L13 ANSWER 27 OF 29 JAPIO COPYRIGHT 2001 JPO

1987-000417 JAPIO ACCESSION NUMBER:

COMPOSITION FOR ORAL CAVITY APPLICATION TITLE:

ICHIKAWA HIROMICHI; KIYOSHIGE TATSUO; KIKUCHI **INVENTOR:**

YASUO

(CO 000676) PATENT ASSIGNEE(S): LION CORP, JΡ

PATENT INFORMATION:

KIND DATE ERA MAIN IPC PATENT NO _____ (4) A61K007-16 JP 62000417 A 19870106 Showa

JP

APPLICATION INFORMATION

19850625 ST19N FORMAT: JP1985-138111 ORIGINAL: JP60138111 Showa

PATENT ABSTRACTS OF JAPAN, Unexamined SOURCE:

Applications, Section: C, Sect. No. 425, Vol.

11, No. 167, P. 128 (19870528)

AN 1987-000417 **JAPIO**

PURPOSE: To provide the titled composition effective to suppress the AB fixation of pathogenic bacteria of periodontosis in oral cavity and prevent the periodontosis, by using the whole cell, pilus or capsule of the pathogenic bacteria of periodontosis as an antigen, immunizing a mammal with the antigen and using the obtained antibody in combination with a nonionic surfactant as active components.

CONSTITUTION: The objective composition contains (A) an antibody produced by immunizing a mammal with an antigen comprising the whole cell, pilus or capsule of the pathogenic bacteria of periodontosis (e.g. Bacterroides gingivalis) (preferably compounded with an antiserum or mild containing said antibody) and (B) preferably 0.1-5wt% nonionic surfactant (preferably fatty acid alkanolamide, sucrose fatty acid ester, polyoxyethylene sorbitan fatty acid ester, etc.).

L13 ANSWER 28 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

1985-297784 [48] ACCESSION NUMBER:

DOC. NO. CPI: C1985-128762

TITLE: Stabilised compsn. contq. basic protein or poly peptide - with a modified gelatin as stabiliser,

esp. for an interferon obtd. by recombinant DNA

WPIDS

technology.

DERWENT CLASS: B04 D16 INVENTOR(S): TERANO, Y

(SUNR) SUNTORY LTD PATENT ASSIGNEE(S):

COUNTRY COUNT: 13

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG ______

A 19851127 (198548) * EN EP 162332

R: AT BE CH DE FR GB IT LI LU NL SE

JP 60228422 A 19851113 (198601) US 4659570 A 19870421 (198718)

EP 162332 B 19890719 (198929) EN
R: AT BE CH DE FR GB IT LI LU NL SE
DE 3571552 G 19890824 (198935)
JP 04081573 B 19921224 (199304) 9

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
EP 162332	A	EP 1985-105092	19850426
JP 60228422	A	JP 1984-84990	19840426
US 4659570	A	US 1985-727261	19850425
JP 04081573	В	JP 1984-84990	19840426

FILING DETAILS:

PATENT NO	KIND		PAT	TENT NO
JP 04081573	В	Based o	on JP	60228422

PRIORITY APPLN. INFO: JP 1984-84990 19840426

AN 1985-297784 [48] WPIDS

AB EP 162332 A UPAB: 19930925

Stabilised compsn. contains a physiologically active basic protein (I) or polypeptide (II) together with a modified gelatin (III).

Pref. the compsn. esp. contains an interferon which is a polypeptide consisting of 146 amino acid residues arranged in the same sequence as in human gamma-interferon or a polypeptide partly deficient of its C terminal and having the activity of human gamma-interferon (IFN-gamma).

(III) is obtd. by (a) decomposing a physically modified gelatin and forming urea bridges by treatment with a diisocyanate; or (b) decomposing the physically modified gelatin and succinylating the prod. with succinic anhydride; or (c) condensing the physically modified gelatin with glyoxal, then oxidising the prod. with H2O2. The physically modified gelatin is easily water soluble and is obtd. by spray-drying or freeze-drying a gelatin or drying by radiofrequency induction heating. The compsn. may also contain an antiviral non-ionic surfactant, anionic surfactant, a human serum albumin, a sugar, etc.

An isotonic agent such as an inorganic salt may also be included.

USE/ADVANTAGE - (I) and (II) are esp.obtd. by cultivation of a micro-organism transformed by a recombinant DNA, and they include interferons, interleukin 2, lysozyme and antibody complement. (III) is more effective as a stabiliser than prior materials such as human serum albumin.

0/0

ABEQ EP 162332 B UPAB: 19930925

A prepn of stabilised polypeptide having gamma-interferon activity which comprises a polypeptide having gamma-interferon activity and a modified gelatin obtd by (1) hydrolsying a physically modified gelatin and forming urea bridges by treatment with a diisocyanate, or (2) hydrolysing said physically modified gelatin and succinylating the hydrolysis product with succinic anhydride, or (2) condensing said physically modified gelatin with glyoxal and oxidising the condensation product with hydrogen peroxide, said physically modified gelatin being easily water soluble and obtd by spray-drying or freeze-drying a telatin, or

drying the same by radiofrequency induction heating.

ABEO JP 92081573 B UPAB: 19930925

Stabilised prepn. of physiologically active substance comprising a basic protein or a polypeptide and a modified gelatin is claimed. The basic protein or a polypeptide is obtd. from a cultivated prod. of recombined micro-organism made by recombination DNA technology, and the physiologically active substance is an interferon. The interferon is a polypeptide consisting of 146 aminoacid radicals of the same sequence as in human-gamma interferon, or a polypeptide which lacks part or its C terminal and has human gamma-interferon activity.

USE - Method is esp. effective for stabilised prepn. of INF-gamma. (J60228422-A)

4659570 A UPAB: 19930925 ABEO US

A poplypeptide (PP) having gamma-interferon (GI) activity is stabilised by a gelatin obtd. by (A) decomposing a physically modified gelatin (PMG) and forming urea bridges by treatment with a diisocyanate, (B) decomposing a PMG and succinylating it with succinic anhydride or (C) condensing the PMG with glyoxal and oxidising the condensation product with H2O2. The PMG is readily soluble in water and is obtd. by spray or freeze drying a gelatin or drying it by radio-frequency induction heating.

The PP is pref. obtd from a culture of a microorganism transformed by a recombinant DNA, and either has the correct amino acid sequence of human GI or has the activity of human GI. The stabilised PP also contains an antiviral non-ionic surfactant, an anionic surfactant, a human serum albumin and a sugar as well as an isotonic agent, e.g. an inorganid salt.

ADVANTAGE - The PP can be stored for prolonged period at room temp; the stabiliser does not affect the activity of the PP.

L13 ANSWER 29 OF 29 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER:

1982-86226E [41] WPIDS

TITLE:

Homogeneous bovine, horse and sheep erythrocyte glyco proteins - useful in sensitive diagnostic

haemagglutination assays.

DERWENT CLASS:

B04 C03 D16 S03

INVENTOR(S):

FLETCHER, M A

PATENT ASSIGNEE(S):

(UYMI-N) UNIV MIAMI

COUNTRY COUNT:

12

PATENT INFORMATION:

PAT	TENT NO	KIND	DATE	WEEK	LA	PG
EP				(198241)*	EN	39
			FR GB LI N			
JΡ	57206694	Α	19821218	(198305)		
US	4460694	Α	19840717	(198431)		
US	4525459	Α	19850625	(198528)		
CA	1198051	Α	19851217	(198604)		
ΕP	61912	В	19870930	(198739)	EN	
	R: BE CH	DE	FR GB LI N	NL SE		
DE	3277412	G	19871105	(198745)		
JP	05194599	Α	19930803	(199335)		16
JP	07035397	B2	19950419	(199520)		15

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
EP 61912 US 4460694	A A	EP 1982-301605 US 1982-356348	19820326 19820309
US 4525459	A	US 1982-343235	19820127
JP 05194599	A Div ex	JP 1982-48786 JP 1991-216713	19820326 19820326
JP 07035397	B2	JP 1982-48786	19820326

FILING DETAILS:

PATENT NO F	KIND	PATENT NO
JP 07035397	B2 Based on	JP 57206694

PRIORITY APPLN. INFO: US 1981-247934 19810326; US 1982-343235 19820127; US 1982-356348 19820309; US 1983-343235 19830127

AN 1982-86226E [41] WPIDS

AB EP 61912 A UPAB: 19930915

Homogeneous borine glycoprotein (I) from bovine erythrocytes is new. It has an amino acid compsn., in mole %, of aspartic acid 7.2, threonine 8, serine 7.2, glutamic acid 16.5, proline 12.9, glycine 8.9, alanine 5.6, valine 5.4, methionine 1.2, isoleucine 6.4, leucine 9.2, tyrosine 0.9, phenylalanine 2.8, histidine 1.3, lysine 1.8 and arginine 4.8. The (I) is free from glycolipids and it contains 25% by wt. of carbohydrate comprising hexose, sialic acid, N-acetylgalactosamine and N-acetyl glucosamine (molar ratio 1.6:1:0.5:1.1) and 75% protein (I) gives a single band on polyacrylamide gel electrophoresis when stained with Coomassie blue or with HIO4 modified Schiff reagent.

New horse (II) and sheep (III) glycoproteins are also new along with a procedure for diagnosine mononucleosis.

With (I)-(III) in the homogeneous forms, there is at least a 10-fold increase in sensitivity of the diagnostic haemagglutination tests etc. in which they can be used, compared with the use of the prior crude erythrocyte prepns. (II) and (III) interact with peripheral blood lymphocytes to form E rosettes in vitro and so are useful for enumerating rosetting lymphocytes. (I)-(III) are esp. useful in rapid detection and quantification of antibody to Epstein-Barr virus.

ABEQ US 4460694 A UPAB: 19930915

Bovine glycoprotein having approx. amino acid compsn. (mol.%): 7.2 threonine, 7.2 serine, 16.5 glutamic acid, 12.9 proline, 8.9 glycine, 5.6 alanine, 5.4 valine, 1.2 methionine, 6.4 isoleucine, 9.2 leucine, 0.9 tyrosine, 2.8 phenylalanine, 1.3 histidine, 1.8 lysine and 4.8 arginine is new.

USE/ADVANTAGE - The bovine glycoprotein can be labelled with a radioisotope, enzyme, chromophore etc. and used in determn. and detection of heterophile antobody of human infectious mononucleosis.

New horse erythrocyte glycoprotein has amino acid compsn. of 8.1 (mol.)% aspartic acid, 10.6% threonine, 10.8% serine, 9.4% glutamic acid, 12.3% proline, 9.2% glycine, 11.3% alanine, 4.4% valine, 0.8% methionine, 3.5% isoleucine, 8.2% leucine, 1.1% tyrosine, 2.9% phenylalanine, 1.2% histidine, 1.3% lysine and 4.8% arginine.

New sheep erythrocytes glycoprotein has amino acid compsn. of 5.6 (mole.)% aspartic acid, 8.1% threonine, 12.9% serine, 13.0% glutamic acid, 11.6% proline, 7.7% glycine. 9.6% alanine, 6.2% valine, 0.5% methionine, 4.6% isoleucine, 8.3% leucine, 4.6% tyrosine, 1.2% phenylalanine, 1.6% histidine, 3.2% lysine, 4.0% arginine and 0.3% tryptophan.

USE - For detection of infectious mononucleosis heterophile antibodies and for prepn. of a stable standardisable reagent for counting rosetting lymphocytes.

ABEQ EP 61912 B UPAB: 19930915

A process for preparing a bovine, horse or sheep erythrocyte glycoprotein, which comprises the steps of: (a) uniformly suspending dried, ground, hemoglobin-free stroma from the appropriate erythrocytes in anhydrous acetone; (b) refluxing for from about 1 to about 6 hours, filtering and drying the residue; (c) suspending said dried residue in 100% anhydrous ethanol; (d) refluxing for from 1 to about 6 hours, filtering and drying the residue; (e) suspending the dried residue from step (d) in aqueous ethanol of from about 50% to about 80% strength and repeating step (b); (f) dissolving the residue from step (e) in water and adding 90% aqueous ethanol, followed by incubating on ice, until crystallisation occurs, centrifuging and dialysing the solid layer against a low pH, low ionic strength buffer; (g) passing the solid from step (f) through a cation exchange resin on a chromatographic column; (h) collecting the sialic acid containing fractions from the column and drying them; (i) treating the collected fractions from step (h) by extraction with a known lipid solvent, centrifuging, collecting the aqueous layer and drying it; (j) repeating step (i) on the product of that step, using a different lipid solvent; and (k) recovering the product of step (j) in lyophilised form; characterised in that complex glycolipid is removed from the product of step (k) by: (1) dissolving the product of step (k) in a low ionic strength buffer containing about 1% neutral detergent; (m) loading the solution from step (1) on an anion exchange chromatographic column; (n) washing the column thoroughly with low ionic strength buffer; (o) eluting the column with aqueous buffer to high salt concentration; and (p) dialysing the product of step (o) against water and recovering the product in freeze dried form.

=> fil hom FILE 'HOME' ENTERED AT 12:42:29 ON 04 DEC 2001

	PRINT PRINCIP	STRY ENTERED AT 12:10:49 ON 04 DEC 2001 E POLYSORBATE/CN
		E POLYSORBATES/CN
L1	9	S POLYSORBATE ?/CN E "POLYOXYETHYLENE-POLYOXYPROPYLENE POLYMER"/CN 5
L2		S E3
L3	10	S L1 OR L2
L5	38	E TWEEN/CN 5 S TWEEN ?/CN
L7	22	S (GLUCOSE OR MANNOSE OR GALACTOSE OR FRUCTOSE OR SORBOSE E "N-METHYLGLUCOSAMINE"/CN 5
L8 L9		S E3 S L7 OR L8
		KAMAR
L1	FILE CARL 9	US! ENTERED AT 12:21:41 ON 04 DEC 2001 SEA FILE=REGISTRY ABB=ON PLU=ON POLYSORBATE ?/CN
L2	1	SEA FILE=REGISTRY ABB=ON PLU=ON "POLYOXYETHYLENE-POLYOX YPROPYLENE POLYMER"/CN
L3		SEA FILE=REGISTRY ABB=ON PLU=ON L1 OR L2
L4	1260	SEA FILE=CAPLUS ABB=ON PLU=ON (LYOPHILIZ? OR LYOPHILIS? OR FREEZ?(W)(DRY? OR DRIED)) AND (MOAB OR MAB OR ANTIBOD?)
L5		SEA FILE=REGISTRY ABB=ON PLU=ON TWEEN ?/CN
L6	80	SEA FILE=CAPLUS ABB=ON PLU=ON L4 AND (L3 OR L5 OR TWEEN OR DETERGENT OR SURFACTANT OR SURFAC? (2A) ACTIVE OR POLYSORBATE OR POLY SORBATE OR (POLYOXYETHYLENE OR POLY(W) (OXYETHYLENE OR OXY ETHYLENE) OR POLYOXY ETHYLENE) (3A) (POLYOXYPROPYLENE OR POLY(W) (OXYPROPYLENE OR OXY PROPYLENE) OR POLYOXY PROPYLENE))
L7	22	SEA FILE=REGISTRY ABB=ON PLU=ON (GLUCOSE OR MANNOSE OR GALACTOSE OR FRUCTOSE OR SORBOSE OR SUCROSE OR LACTOSE OR MALTOSE OR CELLOBIOSE OR GENTIOBIOSE OR ISOMALTOSE OR TREHALOSE OR RAFFINOSE OR GLUCOSAMINE OR "N-METHYL-GLUCOS AMINE" OR GALACTOSAMINE OR NEURAMINIC ACID)/CN
L8	1	SEA FILE=REGISTRY ABB=ON PLU=ON N-METHYLGLUCOSAMINE/CN
L9 L10		SEA FILE=REGISTRY ABB=ON PLU=ON L7 OR L8 SEA FILE=CAPLUS ABB=ON PLU=ON L9 OR SUGAR OR GLUCOSE OR MANNOSE OR GALACTOSE OR FRUCTOSE OR SORBOSE OR SUCROSE OR LACTOSE OR MALTOSE OR CELLOBIOSE OR GENTIOBIOS E OR ISOMALTOSE OR TREHALOSE OR RAFFINOSE OR GLUCOSAMINE OR METHYLGLUCOSAMINE OR GALACTOSAMINE OR NEURAMINIC
L11	37	SEA FILE=CAPLUS ABB=ON PLU=ON L6 AND (L10 OR ?SACCHARID E?)
	ANSWER 1 O SION NUMBE	R: 2001:800884 CAPLUS The effects of Tween 20 and sucrose on the stability of anti-L-selectin during lyophilization
AUTHC	PR(S):	<pre>and reconstitution Jones, Latoya S.; Randolph, Theodore W.; Kohnert, Ulrich; Papadimitriou, Apollon; Winter, G.; Hagmann, Marie-Luise; Manning, Mark C.; Carpenter, John F.</pre>
CORPC	RATE SOURC	E: School of Pharmacy, University of Colorado

Searcher :

Shears

308-4994

Wiley-Liss, Inc.

We have chosen an anti-L-selectin antibody as a model

Journal

English

SOURCE:

PUBLISHER:
DOCUMENT TYPE:

LANGUAGE:

Health Sciences Center, Denver, CO, 80262, USA

308-4994

Shears

J. Pharm. Sci. (2001), 90(10), 1466-1477 CODEN: JPMSAE; ISSN: 0022-3549

protein to investigate the effects of sucrose and/or Tween 20 on protein stability during lyophilization and reconstitution. Native anti-L-selectin secondary structure is substantially retained during lyophilization in the presence of sucrose (1 or 0.125%). However, aggregation of the protein during reconstitution of lyophilized protein powders prepd. without sucrose is not reduced by the presence of sucrose in the reconstitution medium. Aggregate formation upon reconstitution is completely inhibited by freeze drying the protein with sucrose and reconstituting with a 0.1% Tween 20 soln. Tween 20 (0.1%) also partially inhibits loss of native anti-L-selectin secondary structure during lyophilization. However, upon reconstitution the formulations lyophilized with Tween 20 contain the highest levels of aggregates. The presence of Tween in only the reconstitution soln. appears to inhibit the transition from dimers to higher order oligomers. Potential mechanism(s) for the Tween 20 effects were investigated. However, no evidence of thermodn. stabilization of anti-L-selectin conformation (e.g., by Tween 20 binding) could be detected. REFERENCE COUNT: 39 (1) Allison, S; Arch Biochem Biophys 1998, V358, REFERENCE(S): P171 CAPLUS (2) Allison, S; Arch Biochem Biophys 1999, V365, P289 CAPLUS (3) Allison, S; Biophys J 1996, V71, P2022 CAPLUS (5) Arakawa, T; Adv Drug Del Rev 1993, V10, P1 CAPLUS (7) Bam, N; Biotechnol Prog 1996, V12, P801 CAPLUS ALL CITATIONS AVAILABLE IN THE RE FORMAT L11 ANSWER 2 OF 37 CAPLUS COPYRIGHT 2001 ACS 2001:783286 CAPLUS ACCESSION NUMBER: Effect of moisture on the stability of a TITLE: lyophilized humanized monoclonal antibody formulation Breen, E. D.; Curley, J. G.; Overcashier, D. E.; AUTHOR(S): Hsu, C. C.; Shire, S. J. Pharmaceutical Research and Development, CORPORATE SOURCE: Genentech, Inc., South San Francisco, CA, 94080, USA Pharm. Res. (2001), 18(9), 1345-1353 SOURCE: CODEN: PHREEB; ISSN: 0724-8741 Kluwer Academic/Plenum Publishers PUBLISHER: DOCUMENT TYPE: Journal English LANGUAGE: Purpose. To det. the effect of moisture and the role of the glass AB transition temp. (Tg) on the stability of a high concn.,

Searcher:

lyophilized, monoclonal antibody. Methods. A humanized monoclonal antibody was lyophilized in a sucrose/histidine/polysorbate 20 formulation. Residual moistures were from 1 to 8%. Tg values were measured by modulated DSC. Vials were stored at temps. from 5 to 50.degree.C for 6 or 12 mo. Aggregation was monitored by size exclusion chromatog. and Asp isomerization by hydrophobic interaction chromatog. Changes in secondary structure were monitored by Fourier transform IR (FTIR). Results. Tg values varied from 80.degree.C at 1% moisture to 25.degree.C at 8% moisture. There was no cake collapse and were no differences in the secondary structure by FTIR. All formulations were stable at 5.degree.C. High moisture cakes had higher aggregation rates than drier samples if stored above their Tg values. Intermediate moisture vials were more stable to aggregation than dry vials. High moisture samples had increased rates of Asp isomerization at elevated temps. both above and below their Tg values. Chem. and phys. degrdn. pathways followed Arrhenius kinetics during storage in the glassy state. Only Asp isomerization followed the Arrhenius model above the Tg value. Both chem. and phys. stability at T .gtoreq. Tg were fitted to Williams-Landel-Ferry (WLF) kinetics. The WLF consts. were dependent on the nature of the degrdn. system and were not characteristic of the solid system. Conclusion. High moisture levels decreased chem. stability of the formulation regardless of whether the protein was in a glassy or rubbery state. In contrast, phys. stability was not compromised,

and may even be enhanced, by increasing residual moisture if storage

is below the Tg value. REFERENCE COUNT:

REFERENCE(S):

- (2) Bell, L; J Pharm Sci 1995, V84, P707 CAPLUS
- (3) Boye, J; J Agricul Food Chem 1997, V45, P1116 CAPLUS
- (4) Cacia, J; Biochemistry 1996, V35, P1897 CAPLUS
- (5) Carpenter, J; Dev Biol Stand 1992, V74, P225 CAPLUS
- (6) Chang, B; Arch Biochem Biophys 1996, V331, P249 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 3 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: DOCUMENT NUMBER:

2001:780648 CAPLUS

TITLE:

Polymer-based injectable sustained release pharmaceutical compositions for peptide and

protein drugs

135:335147

INVENTOR(S):

Lee, Hee-yong; Lee, Hye-suk; Kim, Jung-soo; Kim,

Sang-beom; Lee, Ji-suk; Choi, Ho-il; Chang,

. Seung-gu

PATENT ASSIGNEE(S):

SOURCE:

Peptron Inc., S. Korea PCT Int. Appl., 37 pp. CODEN: PIXXD2

Patent

DOCUMENT TYPE: LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

KIND DATE PATENT NO.

APPLICATION NO. DATE

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WO 2001078687
                         Α1
                               20011025
                                                WO 2001-KR462
              AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KZ, LC, LK, LR,
              LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ,
              ΜT
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
              TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD,
              TG
PRIORITY APPLN. INFO.:
                                            KR 2000-20484
                                                               A 20000418
                                                               A 20000824
                                            KR 2000-49344
     Controlled and sustained release injectable pharmaceutical compns.
     for a biopharmaceutical, such as peptides and proteins are
     described. Processes for prepn. of an injectable sustained release
     compn. comprises (i) a step of prepg. biodegradable porous
     microspheres having accessible ionic functional groups, (ii) a step
     of encapsulating a biopharmaceutical into the microspheres through
     ionic interaction by suspending or equilibrating the microspheres in
     a soln. contg. the biopharmaceutical, and (iii) a step of recovering
     and freeze-drying the biopharmaceutical-
     incorporated microspheres. For example, microspheres were prepd. by
     water/oil/water double emulsion solvent evapn. method using a
     hydrophilic 50:50 PLGA polymer (RG 502H), which contains free
     carboxy end groups. Deionized water (800 mL) was added to 1 g of
     PLGA polymer dissolved in 2 mL of methylene chloride and emulsified
     by sonication for 30 s using a probe type ultrasonic generator.
     This primary emulsion was dispersed into 200 mL of deionized water
     contg. 0.5% polyvinyl alc. (wt./vol.) in a vessel which connected to
     a const. temp. controller and mixed well by stirring for 15 min at
     2500 rpm, 25.degree. using a mixer. After mixing for another 15 min
     at 1500 rpm, 25.degree., temp. of continuous phase was increased to
     40.degree. to evap. methylene chloride. After 1 h stirring at
     40.degree., 1500 rpm, temp. was decreased to 25.degree.. The
     hardened microspheres were collected by centrifugation and washed
     twice with 200 mL of deionized water, and then freeze-
     dried. The microspheres obtained were used for
     incorporation of protein drugs, i.e., ovalbumin, bovine serum
     albumin, human growth hormone, RNase A, or lysozyme through ionic
     interaction by simply soaking and equilibrating the microspheres
     into a buffer soln. having an appropriate concn. of protein.
REFERENCE COUNT:
                            (1) Bodmer; In J Controlled Release 1992,
REFERENCE(S):
                                V211-3, P129
                            (2) Syntex Inc; US 5470582 1995 CAPLUS
                       CAPLUS COPYRIGHT 2001 ACS
L11 ANSWER 4 OF 37
                            2001:50923 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                            134:114834
TITLE:
                            Preparation of antibody or antigen
                            spheres for diagnostic tests
                           Goertz, Susan; Hemmes, Paul
INVENTOR(S):
                            Spectral Diagnostics, Inc., Can.
PATENT ASSIGNEE(S):
SOURCE:
                            PCT Int. Appl., 34 pp.
                            CODEN: PIXXD2
DOCUMENT TYPE:
                            Patent
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AB

308-4994 Searcher : Shears

LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: KIND DATE APPLICATION NO. PATENT NO. ---------------____ A2 20010118 WO 2000-IB967 20000714 WO 2001004633 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TMRW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG US 1999-353191 A 19990714 US 2000-616018 A 20000713 PRIORITY APPLN. INFO.: US 2000-616018 The invention relates to surfactant-free reagent spheres AB useful for biol. tests particularly those involving antigen/ antibody reactions. The prepd. antibody or antigen spheres are used for detection of microorganism, peptidoglycan, lipopolysaccharide, cytokine, interleukin, tumor necrosis factor, drug abuse, or therapeutic agent; and for diagnosis of infection, sepsis, etc. diseases. 50-99-7, Glucose, biological studies ΙT RL: ANT (Analyte); BSU (Biological study, unclassified); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses) (prepn. of antibody or antigen spheres for diagnostic tests) 99-20-7, Trehalose ΙT RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses) (prepn. of antibody or antigen spheres for diagnostic tests) L11 ANSWER 5 OF 37 CAPLUS COPYRIGHT 2001 ACS 2001:27412 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 134:105824 CD18- or CD20-binding antibody TITLE: formulation Lam, Xanthe M.; Oeswein, James Q.; INVENTOR(S): Ongpipattanakul, Boonsri; Shahrokh, Zahra; Wang, Sharon X.; Weissburg, Robert P.; Wong, Rita L. Genentech, Inc., USA PATENT ASSIGNEE(S): SOURCE: U.S., 56 pp. CODEN: USXXAM DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6171586	B1	20010109	US 1998-97171	19980612
PRIORITY APPLN. INFO.	:		US 1997-53087 P	19970613

09/308223 A stable aq. pharmaceutical formulation comprising a therapeutically AΒ effective amt. of an antibody not subjected to prior lyophilization, a buffer maintaining the pH in the range from about 4.5 to about 6.0, a surfactant and a polyol is described, along with uses for such a formulation. ΙT 57-50-1, Sucrose, biological studies 99-20-7, Trehalose RL: MOA (Modifier or additive use); PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses) (CD18- or CD20-binding antibody formulation) REFERENCE COUNT: REFERENCE(S): (1) Akers, M; Pharmaceutical Technology 1984, P36 CAPLUS (2) Albelda; FASEB-J 1990, V4(11), P2868 CAPLUS (3) Anderson; US 5736137 1998 CAPLUS (4) Anon; EP 303746 1989 CAPLUS (6) Anon; WO 8909402 1989 CAPLUS ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 6 OF 37 CAPLUS COPYRIGHT 2001 ACS 2000:822526 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 134:9337

Adjuvant optimized for stability and TITLE:

biocompatibility for enhancing humoral and

cellular immune responses

Mueller, Rainer Helmut; Grubhofer, Nikolaus; INVENTOR(S):

Olbrich, Carsten

Gerbu G.m.b.H., Germany; Pharmasol G.m.b.H. PATENT ASSIGNEE(S):

Ger. Offen., 26 pp. SOURCE:

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA:	PATENT NO.				ND	D DATE				PPLI			o.	DATE			
WO	1002 2000 2000	0711	54	A	2	2000 2000 2001	1130		DI	E 20	00-1	0024		2000 2000	0519		
	W:	CR, HR, LS, PT,	CU, HU, LT, RO,	CZ, ID, LU, RU,	DE, IL, LV, SD,	DK, IN, MA, SE,	DM, IS, MD, SG,	DZ, JP, MG, SI,	EE, KE, MK, SK,	ES, KG, MN, SL,	FI, KP, MW, TJ,	GB, KR, MX, TM,	GD, KZ, MZ, TR,	CA, GE, LC, NO, TT, MD,	GH, LK, NZ, TZ,	GM, LR, PL, UA,	
	RW:	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	AT, NL, SN,	PT,	SE,	
WO	2000													2000			
	W:	CR, HR, LS, PT,	CU, HU, LT, RO,	CZ, ID, LU, RU,	DE, IL, LV, SD,	DK, IN, MA, SE,	DM, IS, MD, SG,	DZ, JP, MG, SI,	EE, KE, MK, SK,	ES, KG, MN, SL,	FI, KP, MW, TJ,	GB, KR, MX, TM,	GD, KZ, MZ, TR,	CA, GE, LC, NO, TT, MD,	GH, LK, NZ, TZ,	GM, LR, PL, UA,	

WO 2000-EP4644

W 20000522

ΨМ

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

AU 2000058091 A5 20001212 AU 2000-58091 20000522 PRIORITY APPLN. INFO.: DE 1999-19923256 A1 19990520

AB A title adjuvant is disclosed for injection in combination with an antigen. The adjuvant consists of solid lipid particles or solid lipid mixts. It can be used for manuf. of efficient and biocompatible vaccines for immunization of human and other animals as well as for the prodn. of antibodies. By selection of the particle size, particle charge, and particle surface properties the strength of the immune response can be modulated. The optimized adjuvant can be used in combination with other adjuvants such as mol. adjuvants like GMDP.

IT 9003-11-6, Polyoxyethylene-

polyoxypropylene copolymer 9005-65-6,

Tween 80

RL: PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(adjuvant optimized for stability and biocompatibility for enhancing humoral and cellular immune responses)

L11 ANSWER 7 OF 37 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 2000:513486 CAPLUS

DOCUMENT NUMBER: 133:125304

TITLE: Nonaqueous solutions and suspensions of

macromolecules for pulmonary delivery

INVENTOR(S): Klibanov, Alexander M.

PATENT ASSIGNEE(S): Massachusetts Institute of Technology, USA

SOURCE: PCT Int. Appl., 12 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE
WO 2000042993 A2 20000727 WO 2000-US957 20000114
WO 2000042993 A3 20001130

W: CA, JP

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE

PRIORITY APPLN. INFO.:

US 1999-116860 P 19990122
US 1999-443716 A 19991119

AB Methods and formulations for delivery of macromols., such as proteins, polysaccharides, and nucleic acids, are disclosed, where the macromol. is dissolved or dispersed in a low toxicity org. solvent which can be aerosolized for delivery to a patient's lungs by inhalation. Optionally, appropriate soly. enhancers are also present in the formulations compn.

L11 ANSWER 8 OF 37 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 2000:114380 CAPLUS

DOCUMENT NUMBER: 132:171107

09/308223 TITLE: Lyophilized imaging agent formulation comprising a chemotactic peptide Corbo, Diane C.; Link, Mary Jean M.; Williams, INVENTOR(S): N. Adeyinka; Tomsho, Michelle L.; Bornstein, Michael; Solomon, Howard F.; Larsen, Scott K.; Suddith, Robert L. PATENT ASSIGNEE(S): Ortho Pharmaceutical Corp., USA; Johnson-Matthey Inc. U.S., 25 pp., Division of U.S. Ser. No. 271,818, SOURCE: abandoned. CODEN: USXXAM DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: KIND DATE APPLICATION NO. DATE PATENT NO. ____ -----_____ _____ _____

US 6024938 A 20000215 US 1997-997894 19971224 ZA 9505642 A 19970506 ZA 1995-5642 19950706 PRIORITY APPLN. INFO.: US 1994-271818 B3 19940707

A lyophilized imaging agent formulation comprises a targeting mol. such as antibody or chemotactic peptide, a linker such as diethylenetriaminepentaacetic acid (DTPA) or succinimidyl 6-hydrazinium nicotinate-HCl (SHNH), drying protectant such as mannitol, maltose or tricine, and excipient such as Polysorbate 80, in citrate buffer. The formulations are lyophilized and may be stored for extended periods of time. Following reconstitution with a diluent, the formulations are administered to a subject for scintigraphic imaging or therapeutic use. Also contemplated is a kit comprising a 2-vial system wherein a first vial comprises a lyophilized formulation of imaging agent in the form of a lyophilized cake, and a second vial comprises a carrier or diluent. The DTPA-IgG imaging agent was lyophilized in accordance with the following protocol. One-half (0.5) mL of a 4 mg/mL imaging agent formulation (DTPA-IgG) contg. saline 0.9, maltose 5, and Polysorbate-80 0.04% and pH 4.5 citrate buffer (80 mM) was lyophilized. The formulation was placed in a glass vial and the vial held in the lyophilizer at 5.degree. for one-half hour, after which time the temp. was ramped-down over 9 h to -50.degree.. The lyophilized vial was reconstituted with 2.0 mL 0.9% saline and the final formulation contained imaging agent 1 mg/mL in 20 mM citrate buffer, 1.25% maltose, 0.9% saline and 0.01% Polysorbate-80 at pH 4.5.

IT 69-79-4, Maltose 9005-65-6,

Polysorbate 80

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (lyophilized imaging agent formulation comprising

chemotactic peptide)

REFERENCE COUNT:

OUNT: 45

REFERENCE(S): (1) Abrams, M; J Nucl Me

- (1) Abrams, M; J Nucl Med 1990, V31(12), P2022 CAPLUS
- (2) Anon; EP 0314317 A1 1989 CAPLUS (3) Anon; WO 8911297 1989 CAPLUS
- (4) Anon; WO 9104056 1991 CAPLUS
- (5) Borrebaeck, C; J Immunol Meth 1989, V123, P157 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 9 OF 37 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1999:610563 CAPLUS

DOCUMENT NUMBER: 131:241966

TITLE: Stabilization of hepatitis C virus antigen-sensitized latex reagent

INVENTOR(S): Taiheiraku, Yoshihiro; Ifuku, Yasuo; Miyoshi,

Kinya; Washisu, Masayoshi

PATENT ASSIGNEE(S): Mitsubishi Chemical Industries Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 11258241 A2 19990924 JP 1998-56573 19980309

AB Latex particles sensitized with antigen or antibody is

freeze-dried for stable long-term storage.

Dispersing agent, stabilizer, carbohydrate, surfactant and antioxidant are added for stabilization of the immunoassay reagent. A test kit comprising such latex particles sensitized with hepatitis C virus antigen (esp. C25 antigen) is provided for detecting HCV-specific antibody in blood plasma or serum samples.

IT 57-50-1, Sucrose, analysis

RL: ARU (Analytical role, unclassified); MOA (Modifier or additive use); ANST (Analytical study); USES (Uses)

(stabilization of hepatitis C virus antigen-sensitized latex reagent)

L11 ANSWER 10 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1999:581581 CAPLUS

DOCUMENT NUMBER: 132:26728

TITLE: Niosomes as a novel peroral vaccine delivery

system

AUTHOR(S): Rentel, C. -O.; Bouwstra, J. A.; Naisbett, B.;

Junginger, H. E.

CORPORATE SOURCE: Division of Pharmaceutical Technology,

Leiden/Amsterdam Center for Drug Research,

Leiden, 2300 RA, Neth.

SOURCE: Int. J. Pharm. (1999), 186(2), 161-167

CODEN: IJPHDE; ISSN: 0378-5173

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

AB The feasibility to develop a peroral vaccine delivery system based

on nonionic surfactant vesicles (niosomes) was evaluated using BALB/c mice. Ovalbumin was encapsulated in various

lyophilized niosome prepns. consisting of sucrose

esters, cholesterol and dicetyl phosphate. Two different formulations were compared in this study. The specific

antibody titers within serum, saliva and intestinal washings
were monitored by ELISA on days 7, 14, 21 and 28 after intragastric
administration. Only encapsulation of ovalbumin into Wasag 7 (70%)

stearate sucrose ester, 30% palmitate sucrose

ester (40% mono-, 60% di/tri-ester)) niosomes resulted in a significant increase in antibody titers. Administration of ovalbumin and empty niosomes did not exert a similar effect, neither did administration of any control formulation. In contrast to ovalbumin loaded Wasag 7 niosomes, application of the more hydrophilic Wasa 15 (30% stearate sucrose ester, 70% palmitate sucrose ester (70% mono-, 30% di/tri-ester)) niosome prepns. did not result in an increase in antibody titers.

TΨ 57-50-1D, Sucrose, esters

RL: PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES

(niosomes as peroral vaccine delivery systems)

REFERENCE COUNT:

14 REFERENCE(S):

- (1) Alexander, J; WO 93/19781 1993 CAPLUS
- (2) Eldridge, J; J Control Release 1990, V11, P205 CAPLUS
- (3) Elson, C; J Immunol Methods 1984, V67, P101 CAPLUS
- (4) Engvall, E; Methods Enzymol 1980, V70, P419 CAPLUS
- (5) Fujii, Y; Immunol Lett 1993, V36, P65 CAPLUS ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2001 ACS L11 ANSWER 11 OF 37

ACCESSION NUMBER:

1999:354978 CAPLUS

DOCUMENT NUMBER:

131:134516

TITLE:

Lyophilization of Protein Formulations in Vials: Investigation of the Relationship between Resistance to Vapor Flow during Primary Drying and Small-Scale Product Collapse

AUTHOR(S):

Overcashier, David E.; Patapoff, Thomas W.; Hsu,

Chung C.

CORPORATE SOURCE:

Department of Pharmaceutical Research and Development, Genentech Inc., South San

Francisco, CA, 94080, USA

SOURCE:

J. Pharm. Sci. (1999), 88(7), 688-695

CODEN: JPMSAE; ISSN: 0022-3549

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

During the lyophilization process, formulations contg. protein, bulking agent, or lyoprotectant form a dry product layer that can affect the transport of sublimed water vapor. We carried out an investigation of the primary drying segment of lyophilization to evaluate the relationship between the resistance to water vapor flow through the dried layer and the microstructure of the dried cake. Recombinant humanized antibody HER2 (rhuMAb HER2) formulated in trehalose was studied, as were protein-free formulations contg. trehalose and sucrose. Sublimation rate and product temp. data were used to compute the resistance to mass transfer. Dried cake structure was examd. by SEM and a novel fluorescence microscopy method. Collapse temps. were detd. by freeze-drying microscopy. Mass transfer resistance was found to decrease with increases in temp. for each material. Resistance also depended on compn., decreasing in the

> 308-4994 Searcher : Shears

formulation series, rhuMAb HER2, trehalose, sucrose. The lyophilized material consisted of porous cakes, with a distinct denser region at the top. Formulation and temp. affected the microstructure of the dried cakes. The formulated trehalose and sucrose were seen, by both microscopy techniques, to possess small (2-20 .mu.m) holes in their platelike structures after lyophilization. The quantity of holes was higher for material dried at higher temp. The collapse temp. (Tc) of a material appeared to play a role in the process, as lower Tc was correlated with lower resistance and a greater extent of holes. Our results are consistent with the theory that lower resistance to water vapor flow in the primary drying stage of lyophilization may be due to small-scale product collapse.

IT 57-50-1, Sucrose, biological studies 99-20-7, Trehalose 9005-64-5,

Polysorbate 20

RL: PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(lyophilization of protein formulations in vials)

REFERENCE COUNT:

20

REFERENCE(S):

- (1) Carpenter, J; Biochemistry 1989, V28, P3916 CAPLUS
- (2) Chang, B; Pharm Res 1995, V12, P831 CAPLUS(3) Crowe, L; Biophys J 1996, V71, P2087 CAPLUS
- (4) Fendly, B; Cancer Res 1990, V50, P1550 CAPLUS
- (6) Hsu, C; J Pharm Sci 1996, V85, P70 CAPLUS ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 12 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

DOCUMENT NUMBER:

1999:344768 CAPLUS

TITLE:

130:357132
Improved method for stabilizing proteins

INVENTOR(S):

Hellerbrand, Klaus; Papadimitriou, Apollon;

Winter, Gerhard

PATENT ASSIGNEE(S):

Roche Diagnostics G.m.b.H., Germany

SOURCE: Eur. Pat. Appl., 11 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA	TENT NO.	KIND	DATE		APPLICATION NO.	DATE		
EP	917879	A2	19990526		EP 1998-121684	19981113		
EΡ	917879	A3	19990609					
					GB, GR, IT, LI, LU,	NL, SE, MC,		
	PT,	IE, SI, L'	T, LV, FI,	RO				
US	6238664	B1	20010529		US 1998-196090	19981119		
ZA	9810650	A	19990524		ZA 1998-10650	19981120		
AU	9894060	A1	19990610		AU 1998-94060	19981120		
ΑU	714264	B2	19991223					
CN	1220270	Α	19990623		CN 1998-122531	19981120		
BR	9805021	A	20000321		BR 1998-5021	19981123		
JP	11240895	A2	19990907		JP 1998-332681	19981124		

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20001030
     JP 3105494
                       B2
PRIORITY APPLN. INFO.:
                                        EP 1997-120528
                                                         A 19971122
                                        EP 1998-102846
                                                         A 19980219
     Formation of protein aggregates in a reconstituted
AΒ
     lyophilizate of a protein prepn. for pharmaceutical use is
     prevented by prepg. an aq. soln. of the protein in .gtoreq.10 mM K
     phosphate buffer (K:Na ratio .gtoreq.10:1), freezing and thawing the
     soln., dispensing the soln. into injectable doses, and
     lyophilizing. The stabilizing effect of K phosphate buffer
     is attributed at least in part to their small pH shift during
     freezing. The protein soln. preferably also contains a nonionic
     detergent (e.g. polysorbate) and a cryoprotectant
     (e.g. a nonreducing sugar).
L11 ANSWER 13 OF 37 CAPLUS COPYRIGHT 2001 ACS
                         1999:7857 CAPLUS
ACCESSION NUMBER:
                         130:57165
DOCUMENT NUMBER:
                         Stabilized antibody formulation
TITLE:
INVENTOR(S):
                         Lam, Xanthe M.; Oeswein, James Q.;
                         Ongpipattanakul, Boonsri; Shahrokh, Zahra; Wang,
                         Sharon X.; Weissburg, Robert P.; Wong, Rita L.
                         Genentech, Inc., USA
PATENT ASSIGNEE(S):
                         PCT Int. Appl., 87 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                     KIND
                            DATE
                                           APPLICATION NO.
                                                            DATE
     PATENT NO.
                                           _____
                                           WO 1998-US12209 19980612
                            19981217
                      A1
     WO 9856418
         W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
             DE, DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP,
             KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK,
             MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL,
             TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ,
             MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
             ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
             CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
                                          AU 1998-82559
                                                            19980612
     AU 9882559
                       A1
                            19981230
                                           EP 1998-932747
                                                            19980612
                            20000517
     EP 999853
                       A1
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,
             PT, IE, FI
                                        US 1997-874897
                                                            19970613
PRIORITY APPLN. INFO.:
                                        WO 1998-US12209
                                                            19980612
     A stable aq. pharmaceutical formulation comprising a therapeutically
AB
     effective amt. of an antibody not subjected to prior
     lyophilization, a buffer maintaining the pH in the range
     from about 4.5 to about 6.0, a surfactant and a polyol is
     described, along with uses for such a formulation.
     57-50-1, Sucrose, biological studies
     99-20-7, Trehalose
     RL: MOA (Modifier or additive use); PEP (Physical, engineering or
     chemical process); THU (Therapeutic use); BIOL (Biological study);
     PROC (Process); USES (Uses)
```

Searcher: Shears 308-4994

(stabilized antibody formulation)

```
REFERENCE COUNT:
                         (1) Abbott Lab; WO 9641164 A 1996 CAPLUS
REFERENCE(S):
                         (2) Cleland; Critical Reviews in Therapeutic
                             Drug Carrier Systems 1993, V10(4), P307
                             CAPLUS
                         (4) Draber, P; Journal of Immunological Methods
                             1995, V181(1), P37 CAPLUS
                         (5) Genentech Inc; WO 9704801 A 1997 CAPLUS
                         (6) Immuno AG; EP 0661060 A 1995 CAPLUS
                         ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 14 OF 37 CAPLUS COPYRIGHT 2001 ACS
                         1998:527193 CAPLUS
ACCESSION NUMBER:
                         129:166193
DOCUMENT NUMBER:
                         Therapeutic treatment and prevention of
TITLE:
                         infections with a bioactive material
                         encapsulated within a biodegradable-
                         biocompatible polymeric matrix
                         Setterstrom, Jean A.; Van Hamont, John E.; Reid,
INVENTOR(S):
                         Robert H.; Jacob, Elliot; Jeyanthi, Ramasubbu;
                         Boedeker, Edgar C.; McQueen, Charles E.; Tice,
                         Thomas R.; Roberts, F. Donald; Friden, Phil
                         United States Dept. of the Army, USA; Van
PATENT ASSIGNEE(S):
                         Hamont, John E.; et al.
SOURCE:
                         PCT Int. Appl., 363 pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
                         10
PATENT INFORMATION:
                                           APPLICATION NO. DATE
                     KIND DATE
     PATENT NO.
                                           _____
     _____
                      ____
                           _____
                            19980730
                                           WO 1998-US1556 19980127
     WO 9832427
                      A1
            AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
             DE, DK, EE, ES, FI, GB, GE, GH, HU, IL, IS, JP, KE, KG, KP,
             KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX,
             NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR,
             TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU,
             TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES,
             FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
             CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
                                           US 1997-789734
                                                            19970127
     US 6309669
                      В1
                            20011030
                                                            19980127
                                           AU 1998-63175
     AU 9863175
                       A1
                            19980818
                                        US 1997-789734 A 19970127
PRIORITY APPLN. INFO.:
                                                         B1 19840316
                                        US 1984-590308
                                        US 1992-867301
                                                         A2 19920410
                                                         A2 19950522
                                        US 1995-446148
                                                         B2 19950522
                                        US 1995-446149
                                                         B2 19960124
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AB Novel burst-free, sustained release biocompatible and biodegradable microcapsules are disclosed which can be programmed to release their active core for variable durations ranging from 1-100 days in an aq. physiol. environment. The microcapsules are comprised of a core of polypeptide or other biol. active agent encapsulated in a matrix of poly(lactide/glycolide) copolymer, which may contain a

> 308-4994 Searcher : Shears

US 1996-590973

WO 1998-US1556

W 19980127

pharmaceutically acceptable adjuvant, as a blend of upcapped free carboxyl end group and end-capped forms ranging in ratios from 100/0 to 1/99.

IT 9001-62-1, Lipase

RL: BPR (Biological process); DEV (Device component use); PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(prevention of infections with a bioactive material encapsulated within a biodegradable-biocompatible polymeric matrix)

IT 9005-64-5, Tween 20 9005-65-6,

Tween 80 9005-67-8, Tween 60

RL: MOA (Modifier or additive use); PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(prevention of infections with a bioactive material encapsulated within a biodegradable-biocompatible polymeric matrix)

L11 ANSWER 15 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1998:351786 CAPLUS

DOCUMENT NUMBER:

129:32337

TITLE:

Stable lyophilized pharmaceutical

substances from monoclonal or polyclonal

antibodies

INVENTOR(S):

Kallmeyer, Georg; Winter, Gerhard; Klessen,

Christian; Woog, Heinrich

PATENT ASSIGNEE(S):

Boehringer Mannheim G.m.b.H., Germany;

Kallmeyer, Georg; Winter, Gerhard; Klessen,

Christian; Woog, Heinrich

SOURCE:

PCT Int. Appl., 39 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:				KIND DATE APPLICATION NO. DATE												
	9822 9822	136		A	2									1997	1119	
	W:	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,
														ΚE,		
		KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,
				-		-								ТJ,		
		TT,	UA,	UG,	US,	UZ,	VN,	YU,	ZW,	AM,	ΑZ,	BY,	KG,	KZ,	MD,	RU,
		TJ,														
	RW:													DK,		
											SE,	BF,	ВJ,	CF,	CG,	CI,
						MR,						.	_			
	8529			A	1	1998	0715		E	P 19	96-1	1848	9	1996	1119	
	R: 9854	DE			_											
									A	0 19	98-5	4841		1997	1119	
	7354								_					4000		
	9710															
ĒΡ	9411															
	R:			CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	PT,
		IE,	FI	_					•				_	4000		
	1244															
BR	9713	521		A		2000	0321		В.	к 19	97-1	3521		1997	1119	

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JP 2001503781
                        T2
                             20010321
                                             JP 1998-523210
                                                               19971119
PRIORITY APPLN. INFO .:
                                          EP 1996-118489
                                                            Α
                                                               19961119
                                                            W 19971119
                                          WO 1997-EP6452
     Lyophilized therapeutic and diagnostic agents prepd. from
AB
     monoclonal or polyclonal antibodies contain a
     sugar or amino sugar as well as an amino acid and
     a surfactant as stabilizers. These lyophilized
     compns. are stable under refrigeration, at room temp., or even at
     .ltoreq.30.degree. for .ltoreq.2 yr, and are stable after
     reconstitution with water for injection for .ltoreq.5 days.
     compn. contg. monoclonal antibody to hepatitis B virus 8.0, sucrose 58.0, arginine 10.0, Tween 20 0.1 mg, phosphate buffer 15 mM, NaCl 30 mM, H3PO4 to pH 6.5, and water
     for injection to 1.0 mL showed satisfactory stability during storage
     for 4 or 13 wk at 50.degree..
     57-50-1, Sucrose, biological studies
ΙT
     69-79-4, Maltose 99-20-7,
     Trehalose 114-04-5, Neuraminic acid
     512-69-6, Raffinose 3329-30-4, N-
     Methylglucosamine 3416-24-8, Glucosamine
     7535-00-4, Galactosamine 9003-11-6,
     Ethylene oxide/propylene oxide copolymer 9005-64-5,
     Tween 20
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (stable lyophilized pharmaceutical compns. from
        monoclonal or polyclonal antibodies)
L11 ANSWER 16 OF 37 CAPLUS COPYRIGHT 2001 ACS
                        1997:394163 CAPLUS
ACCESSION NUMBER:
                          127:23753
DOCUMENT NUMBER:
                          Stabilization of biological materials by drying
TITLE:
                          without freezing
                          Mattern, Markus; Winter, Gerhard
INVENTOR(S):
                          Boehringer Mannheim Gmbh, Germany
PATENT ASSIGNEE(S):
                          Ger. Offen., 32 pp.
SOURCE:
                          CODEN: GWXXBX
                          Patent
DOCUMENT TYPE:
                          German
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                             APPLICATION NO. DATE
                       KIND
                             DATE
     PATENT NO.
                       ____
                             _____
     DE 19539574
                             19970430
                                             DE 1995-19539574 19951025
                        Α1
     WO 9715288
                        A2
                             19970501
                                             WO 1996-EP4627 19961024
                             19970529
     WO 9715288
                       A3
             AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
             DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ,
             LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL,
             PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US,
             UZ, VN, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB,
             GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM
                                             CA 1996-2235243 19961024
                             19970501
     CA 2235243
                        AA
                             19970515
                                             AU 1996-72984
                                                               19961024
     AU 9672984
                        Α1
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AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT,

EP.1996-934811

19961024

19980812

A2

IE, SI, FI

EP 857060

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CN 1205628
                       Α
                             19990120
                                            CN 1996-199329
                                                              19961024
     BR 9611265
                             19990504
                                            BR 1996-11265
                                                              19961024
                       Α
     JP 11513700
                       Т2
                             19991124
                                            JP 1996-516286
                                                              19961024
                             19980625
                                            NO 1998-1868
                                                              19980424
     NO 9801868
                       Α
PRIORITY APPLN. INFO.:
                                         DE 1995-19539574
                                                              19951025
                                         WO 1996-EP4627
                                                              19961024
     A biol., esp. therapeutic, material is stabilized and preserved by
AB
     prepg. a soln. of (1) the material, (2) a carbohydrate or a
     zwitterionic compd. with polar residues, and (3) a zwitterionic
     compd. with nonpolar residues, and drying the soln. at a temp. above its f.p. The process does not involve use of elevated temps., can
     be carried out in conventional lyophilization app., is
     energy efficient, and is more rapid than freeze
             Thus, a soln. contg. maltose 50,
     drving.
     L-phenylalanine 10, L-arginine 10, polysorbate 80 0.1, and
     recombinant human G-CSF 0.35 mg/mL (pH 7.4) was sterilized by
     filtration and 1-mL portions were dispensed into 2-mL vials fitted
     with lyophilization stoppers and dried isothermally at
     20.degree. and reduced pressure for 48 h. The product had a
     residual water content of 1.16% and a glass transition temp. of
     75.degree.. The content of native (monomeric) G-CSF was still
     99.83% after 13 wk storage at 50.degree..
     57-50-1, biological studies
ΙT
     RL: MOA (Modifier or additive use); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (stabilization of biol. materials by drying without freezing)
L11 ANSWER 17 OF 37 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER:
                         1997:306965 CAPLUS
DOCUMENT NUMBER:
                         127:3884
                         MF59 adjuvant enhances antibody
TITLE:
                         responses of infant baboons immunized with
                         Haemophilus influenzae type b and Neisseria
                         meningitidis group C oligosaccharide
                         -CRM197 conjugate vaccine
                         Granoff, Dan M.; McHugh, Yvonne E.; Raff, Howard
AUTHOR(S):
                         V.; Mokatrin, Ahmad S.; Van Nest, Gary A.
                         Chiron Vaccines, Emeryville, CA, 94608, USA
CORPORATE SOURCE:
SOURCE:
                         Infect. Immun. (1997), 65(5), 1710-1715
                         CODEN: INFIBR; ISSN: 0019-9567
PUBLISHER:
                         American Society for Microbiology
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         English
     The ability of the adjuvant MF59 to enhance the immunogenicity of
AR
     polysaccharide-protein conjugate vaccines was investigated
     in infant baboons. MF59 consists of stable droplets (<250 nm) of
     the metabolizable oil squalene and two surfactants,
     polyoxyethylene sorbitan monooleate and sorbitan trioleate, in an
     oil-in-water emulsion. In humans, MF59 is well tolerated and
     enhances the immunogenicity of recombinant protein subunit or
     particle vaccines. Its effect on the immunogenicity of
     polysaccharide-protein conjugate vaccines is unknown.
     Baboons 1-4 mo of age were immunized i.m. with N. meningitidis group
     C and H. influenzae type b (Hib) oligosaccharide-CRM197
     conjugate vaccines. The lyophilized vaccines were
     reconstituted with phosphate-buffered saline (PBS), Al(OH)3 (alum),
     or MF59. Groups of 5 animals each were given 3 injections of the
     resp. formulations, with one injection every 4 wk. Four weeks after
```

each immunization, the MF59 group had up to 7-fold-higher geometric mean anticapsular-antibody titers than the alum group and 5-10-fold higher N. meningitidis group C bactericidal antibody titers. Twenty-one weeks after the 3rd immunization, the MF59 group still showed 5-10-fold-higher anticapsular antibody titers. The antibody responses of the animals given the vaccines reconstituted with PBS were low at all times measured. Both the MF59 and alum groups, but not the PBS group, showed booster antibody responses to unconjugated Hib and N. meningitidis group C polysaccharides, results consistent with induction of memory B cells. Thus, MF59 may be useful for accelerating and augmenting immunity to polysaccharide-protein conjugate vaccines in infants.

L11 ANSWER 18 OF 37 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1997:218669 CAPLUS

DOCUMENT NUMBER: 126:203735

TITLE: Stable isotonic lyophilized protein

formulation

INVENTOR(S): Andya, James; Cleland, Jeffrey L.; Hsu, Chung

C.; Lam, Xanthe M.; Overcashier, David E.; Shire, Steven J.; Yang, Janet Yu-Feng; Wu,

Sylvia Sau-Yan

PATENT ASSIGNEE(S): Genentech, Inc., USA

SOURCE: PCT Int. Appl., 48 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PA	rent 1	NO.		KIND DATE					A	PPLI	CATI	ON N	٥.	DATE		
WO	9704	801		Α.	1	1997	0213		W	0 19	96-U	S122	51	1996	0723	
	W:	AL,	AM,	AT,	AU,	AZ,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CZ,	DE,	DK,
		EE,	ES,	FI,	GB,	GE,	ΗU,	IL,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LK,
		LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,
		RO,	RU,	SD,	SE,	SG										
	RW:	KE,	LS,	MW,	SD,	SZ,	UG,	AT,	BE,	CH,	DE,	DK,	ES,	FI,	FR,	GB,
		GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,
		GN														
US	6267	958		B :	1	2001	0731		U	S 19	96-6	1536	9	1996	0314	
CA	2226	575		A	Ą	1997	0213		C	A 19	96-2	2265	75	1996	0723	
AU	AU 9665992				1	1997	• • • • • • • • • • • • • • • • • • • •					1996	0723			
	AU 716785															
EP	8459															
	R:								GB,	GR,	ΙT,	LI,	LU,	NL,	SE,	MC,
		PT,	ΙE,	SI,	LT,	LV,	FI									
CN	1191	490		Α		1998	0826		С	N 19	96-1	9583	0	1996	0723	
BR	9609	743		Α		1999	0302		В	R 19	96-9	743		1996	0723	
	1151															
	9800													1998		
US	2001	0143	26													
PRIORIT	Y APP	LN.	INFO	.:										1995		
														1996		
														1996		
														1996		
ΛD 7.	-4-1-1	. 1		1			- fa	~~~·	a+ i a	n ic	doc	arih	~~			

AB A stable lyophilized protein formulation is described

which can be reconstituted with a suitable diluent to generate a high protein concn. reconstituted formulation which is suitable for s.c. administration. For example, anti-IgE and anti-HER2 antibody formulations have been prepd. by lyophilizing these antibodies in the presence of a lyoprotectant. The lyophilized mixt. thus formed is reconstituted to a high protein concn. without apparent loss of stability of the protein.

IT 57-50-1, Sucrose, biological studies
99-20-7, Trehalose

RL: MOA (Modifier or additive use); PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(stable isotonic lyophilized protein formulation)

L11 ANSWER 19 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1996:546594 CAPLUS

DOCUMENT NUMBER: 125:242353

TITLE: Preparation of wax beads containing a reagent

using liquid nitrogen for cooling and

solidifying

INVENTOR(S): Kosak, Kenneth M.; Kosak, Matthew K.

PATENT ASSIGNEE(S): USA

SOURCE: U.S., 14 pp. Cont.-in-part of U.S. 5,413,924.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 5

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5550044	Α	19960827	US 1994-257567	19940610
US 5413924	Α	19950509	US 1992-936357	19920827
US 5968729	Α	19991019	US 1998-49707	19980328
PRIORITY APPLN.	INFO.:		US 1992-835758	19920213
			US 1992-936357	19920827
			US 1994-257567	19940610
			US 1997-918374	19970826

Droplets of molten wax or waxy polymer contq. a reagent are dropped AB onto the surface of liq. nitrogen, the droplets remain on the surface until solidified, and the droplets are removed from the surface before they sink into the lig. nitrogen to provide beads contg. the reagent. The reagent can be any material that can be entrapped in the beads and does not undergo excessive inactivation when the beads are melted by heating to release the reagent. Examples of reagents are heat-resistant enzymes, enzyme substrates, metal salts, oligonucleotides, inclusion compds., surfactants, emulsifiers, antioxidants, stabilizers, drugs, antibiotics, antibodies and antigens. An app. for producing the beads contains a plurality of channels through which liq. nitrogen flows from a reservoir. Each channel passes under a dispenser tip from which droplets are formed and released onto the surface of flowing liq. nitrogen. Liq. nitrogen contg. the beads flows from each channel into a pipe and then over a sepn. sieve. The beads can be used in various in vitro chem., biochem. and immunol. reactions including the PCR, where the reagent is released by heating and melting the beads. The beads have all the combined

features for com. use of: (1) spherical shape, (2) uniform, narrow size range (i.e. .ltoreq.5% deviation), (3) free of water contamination, (4) contain an aq. reagent, and (5) can be produced at high speed (i.e. >1000/min).

L11 ANSWER 20 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1996:544101 CAPLUS

DOCUMENT NUMBER:

125:177462

TITLE:

Surface-modified nanoparticles and method of

making and using them

INVENTOR(S):

Levy, Robert J.; Labhasetwar, Vinod; Song,

Cunxian S.

PATENT ASSIGNEE(S):

USA

SOURCE:

PCT Int. Appl., 170 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.				KI	ND	DATE			A	PPLI	CATI	ON NO	ο.	DATE		
	9620 9620					1996 1998			W	0 19	96-U	S476		1996	0104	
0		AL,		AT,	AU,	CA,		CN,	CZ,	DE,	DK,	GB,	HU,	IS,	JP,	KE,
	RW:			SD, NE,		BE,	CH,	DE,	ES,	FR,	GB,	IT,	LU,	NL,	PT,	SE,
		961		A	A	1996 1996						2079 7556		1996 1996		
	AU 9647556 EP 805678			A.	1	1997	1112		E	P 19	96-9	0347	6	1996	0104	
	R:	AT, PT,	•	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,
JP PRIORITY	1051				2	1998	1117		J US 1				_	1996		
								US 1	995-	3898	93		1995			

Biodegradable controlled-release nanoparticles as sustained release AB bioactive agent delivery vehicles include surface modifying agents to target binding of the nanoparticles to tissues or cells of living systems, to enhance nanoparticle sustained release properties, and to protect nanoparticle-incorporated bioactive agents. Unique methods of making small (10 nm to 15 nm, and preferably 20 nm to 35 nm) nanoparticles having a narrow size distribution which can be surface-modified after the nanoparticles are formed is described. Techniques for modifying the surface include a lyophilization technique to produce a phys. adsorbed coating and epoxy-derivatization to functionalize the surface of the nanoparticles to covalently bind mols. of interest. The nanoparticles may also comprise hydroxy-terminated or epoxide-terminated and/or activated multiblock copolymers, having hydrophobic segments which may be polycaprolactone and hydrophilic segments. The nanoparticles are useful for local intravascular administration of smooth muscle inhibitors and antithrombogenic agents as part of interventional cardiac or vascular catheterization such as a balloon angioplasty procedure; direct application to tissues and/or cells for gene therapy, such as the delivery of osteotropic genes or gene segments into bone progenitor cells; or

oral administration in an enteric capsule for delivery of protein/peptide based vaccines.

L11 ANSWER 21 OF 37 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1996:516694 CAPLUS

DOCUMENT NUMBER: 125:151116

TITLE: Vaccine adjuvants comprising a sulfolipid

polysaccharides Hilgers, Luuk

INVENTOR(S): Hilgers, Luuk
PATENT ASSIGNEE(S): Solvay et Cie., Belg.
SOURCE: PCT Int. Appl., 29 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

					KIND DATE					APPLICATION NO. DATE							
		9620													1995	1221	
		W:	AL,	AM,	AU,	BB,	BG,	BR,	ΒŸ,	CA,	CN,	CZ,	EE,	FI,	GE,	HU,	IS,
			JP,	KG,	KP,	KR,	ΚZ,	LK,	LR,	LT,	LV,	MD,	MG,	MK,	MN,	MX,	NO,
			NZ,	PL,	RO,	RU,	SG,	SI,	SK,	ТJ,	TM,	TT,	UA,	US,	UZ,	VN	
		RW:	KE,	LS,	MW,	SD,	SZ,	UG,	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,
										BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,
			ML,	MR,	ΝE,	SN,	TD,	TG									
	ΒE	1008	978		A!	5	1996	1001		В	E 19	94-1	174	,	1994	1227	
	CA	2208	790		A)	£.	1996	0704		C.	A 19	95-2	2087	90	1995	1221	
		2208													1995	1221	
		9643								Α	U 19	96-4	3248		1995	1221	
		7091															
	BR	9510	223		Α		1997	1230		В	R 19	95-1	0223		1995		
		8148								Ē	P 19	95-9	4200	7	1995	1221	
	EΡ	8148	36		В:	L	2001	0314									
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	PT,
				SI,													
	CN	1171	052		Α		1998	0121		C	N 19	95-1	9709	9	1995	1221	
	CN	1175	264		Α		1998	0304		C	ท 19	95-1	9755	7	1995	1221	
		1996															
		2155															
PRIOR	RITY	APP	LN.	INFO	. :					BE 1	994-	1174		Α	1994	1227	
										WO 1	995-	BE11	9	W	1995	1221	
ΔR	Vac	cina	adii	nvant	- 6 0	יורווור	isin	ma .	2111 f	olin	id n	വയ	acch:	arid	0		

Vaccine adjuvants comprising a sulfolipid polysaccharide in combination with an interface-forming constituent are claimed. The invention also provides a method for prepg. a vaccine by emulsifying an aq. soln. of an antigen and a sulfolipid polysaccharide. The adjuvants are stable at high temps., and are at least as effective as convenient adjuvants. Their local toxicity is generally lower than that of conventional adjuvants. Thus, 6.6. g of lauroyl chloride was added to a soln. of 4.5 g inulin in 100 mL of DMF:pyridine (1:1), stirred and incubated for 6 h at 60.degree. and 18 h at room temp. followed by addn. of 0.6 g chlorosulfonic acid in 10 mL of DMF: pyridine and stirring and incubation at 60.degree. and room temp. was repeated. The solvents were then evapd. under reduced pressure and the sulfolipid polysaccharide thus obtained was dialyzed in phosphate buffered saline for 10 days, then lyophilized. An emulsion contg. 1, Tween 80 2, squalene 10, and phosphate

buffered saline q.s. 100% was stable after 115 days at 37.degree. Guinea pig immunized with vaccines contg. inactivated influenza virus and above adjuvant showed significant elevated serum antibody as compared with the controls.

L11 ANSWER 22 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1996:217300 CAPLUS

TITLE: Development of stable lyophilized monoclonal antibody formulations: Effect of excipients on stability.

AUTHOR(S): Bam, Narendra B.; Dal Monte, Paul R.; Duddu,

Sarma P.

CORPORATE SOURCE: Pharmaceutical Development, SmithKline Beecham

Pharmaceuticals, King of Prussia, PA, 19406, USA Book of Abstracts, 211th ACS National Meeting, New Orleans, LA, March 24-28 (1996), BIOT-143.

American Chemical Society: Washington, D. C.

CODEN: 62PIAJ

DOCUMENT TYPE: Conference; Meeting Abstract

LANGUAGE: English

AB In the development of proteins for therapeutic use, long term

aggregation and degrdn. is usually prevented by

lyophilization. The proteins have been traditionally

protected from the stresses of freezing and drying by addn. of

excipients like sugars, polymers and surfactants

to formulations. We present studies on the effect of sugars

(sucrose, mannitol and trehalose) and

surfactants (Tween) and polymers (PEG, Dextran and

PVP) on the stabilization against aggregation of a monoclonal

antibody. Our studies include FTIR spectroscopy,

Calorimetry and traditional bioanal. techniques for protein characterization. Although monoclonal antibodies differ

significantly in structure to commonly studied globular proteins,

the stabilizing effect of sugars like sucrose

and trehalose has been obsd. to be similar to that

reported in literature. The importance of the relationship between the storage temp. of a lyophile and the glass transition temp. (Tg)

will be stressed. We will also present data comparing and

delineating whether the stabilization occurs during lyophilization or during reconstitution of the dried

product.

L11 ANSWER 23 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1995:576784 CAPLUS

DOCUMENT NUMBER:

122:299132

TITLE: INVENTOR(S):

Liposomes containing particulate materials Gregoriadis, Gregory; Antimisiaris, Sophia

George; Gursel, Ihsan

PATENT ASSIGNEE(S):

United Kingdom Secretary of State for Defence,

London, UK

SOURCE:

SOURCE:

PCT Int. Appl., 35 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.

KIND DATE

APPLICATION NO. DATE

Searcher :

Shears

308-4994

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WO 1994-GB2191 19941007
    WO 9509610 A1 19950413
        W: AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, ES, FI, GB, GE, HU, JP, KE, KG, KP, KR, KZ, LK, LT, LU, LV, MD, MG,
             MN, MW, NL, NO, NZ, PL, PT, RO, RU, SD, SE, SI, SK, TJ, TT,
             UA, US, UZ, VN
         RW: KE, MW, SD, SZ, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT,
             LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR,
             NE, SN, TD, TG
                                           CA 1994-2173601 19941007
     CA 2173601
                      AΑ
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                                           AU 1994-77907
                      A1
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                                                            19941007
    AU 687893
                      B2
                            19980305
                     A1
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    EP 722317
                            19960724
     EP 722317
                     B1
                            20011121
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, SE
                            19960821
                                     GB 1996-7359 19941007
     GB 2297907 A1
     GB 2297907
                      B2
                            19971210
                      Α
                                          BR 1994-7769
     BR 9407769
                            19970318
                                                            19941007
                      T2
                            19970331
                                           JP 1994-510702
                                                            19941007
     JP 09503225
                                                           19941007
                      A 19981202
                                           CN 1994-194325
     CN 1200668
                                                           19941007
                     C1
                          20000210
                                           RU 1996-110886
     RU 2145212
                                        RU 1990 115
US 1996-624556
     US 6322809
                     В1
                            20011127
                                                            19960920
                                                       556 19960920
A 19931007
PRIORITY APPLN. INFO.:
                                        GB 1993-20668
                                        WO 1994-GB2191 W 19941007
     A method is provided for the formation of liposomes of 0.1 .mu.m to
AB
     50 .mu.m in diam. having unilamellar or multilamellar structure and
     contg. water-insol. or undissolved particulate materials comprising
     (a) forming liposomes and removing substantially all of any org.
     solvent used in their prepn., (b) freeze-drying
     the liposomes so formed and then (c) rehydrating them in intimate
     admixt. with the particulate material. Preferred encapsulated
     materials are particulate materials, most preferably microorganisms,
     plant or animal cells or water-insol. structures having org.
     solvent-labile biochem. or immunol. activity, but any water-insol.
     particulate may be encapsulated using the method. For example,
     catalysts or drugs that are sparingly sol. may also be so
     incorporated such that slow release into the patient's body may be
     provided while release of detergents included in many
     liposome prepn. protocols may be avoided.
IT
     99-20-7, Trehalose
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (manuf. of liposomes contg. particulate materials)
L11 ANSWER 24 OF 37 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER:
                    1995:386367 CAPLUS
                         122:142605
DOCUMENT NUMBER:
                        Method for preparing liposomes
TITLE:
                        Hsu, Chung C.
INVENTOR(S):
                         Genentech, Inc., USA
PATENT ASSIGNEE(S):
                         PCT Int. Appl., 26 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
                                           APPLICATION NO. DATE
     PATENT NO.
                      KIND
                            DATE
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WO 9501164
                       A1
                            19950112
                                            WO 1994-US7327
                                                             19940629
         W: AU, CA, JP
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT,
                            19950112
                                            CA 1994-2163860
                                                             19940629
     CA 2163860
                       AA
     AU 9472151
                       A1
                            19950124
                                           AU 1994-72151
                                                             19940629
     AU 689786
                       B2
                            19980409
     EP 706374
                       A1
                            19960417
                                            EP 1995-904347
                                                             19940629
     EP 706374
                       В1
                            19971210
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL,
             PT, SE
     JP 08512056
                       T2
                            19961217
                                            JP 1994-503613
                                                             19940629
     US 5653996
                       Α
                            19970805
                                            US 1995-407424
                                                             19950317
                                        US 1993-84933
                                                             19930630
PRIORITY APPLN. INFO .:
                                        WO 1994-US7327
                                                             19940629
     Methods are provided for the prepn. of liposomes by spraying a soln.
AB
     comprising bilayer-forming matters, and optional addnl. mols. onto
     an aq. surface, through a frequency-generated vibrating nozzle
     without added pressure. The liposomes are useful for the delivery
     of therapeutic, diagnostic, and cosmetic agents. This method
     provides an economic and efficient method of prepg. liposomes on a
     large scale. For example, a soln. contg.
     dipalmitoylphosphatidylcholine, palmitoyloleoylphosphatidylglycerol,
     lung surfactant protein C, and palmitic acid in
     isopropanol was sprayed onto a succinate buffer soln. and the
     obtained liposome soln. was ultrafiltrated via tangential flow
     filtration and lyophilized.
L11 ANSWER 25 OF 37 CAPLUS COPYRIGHT 2001 ACS
                         1994:696634 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         121:296634
TITLE:
                         Lyophilized ligand-receptor complexes
                         for assays and sensors
INVENTOR(S):
                         Ligler, Frances S.; Whelan, James P.
PATENT ASSIGNEE(S):
                         United States Dept. of the Navy, USA; U.S. Drug
                         Testing, Inc.
SOURCE:
                         U.S., 14 pp.
                         CODEN: USXXAM
DOCUMENT TYPE:
                         Patent
                         English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
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PATEN	1 T	10.		KII	ND	DATE			A)	PPLI	CATI	ON NC	ο.	DATE		
US 53				A		1994								1993	-	
WO 95	5027	703		A.	1	19950	0126		W(0 199	94-U	S780	6	1994	0715	
W	₹:	AM,	AT,	AU,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CZ,	DE,	DK,	ES,	FI,
		GB,	GE,	HU,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LK,	LT,	LU,	LV,	MD,	MG,
		MN,	MW,	NL,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SI,	SK,	TJ,	TT,
		UA,	UZ,	VN												
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		SN,	TD,	TG												
CA 21	1672	275		A)	Ą	1995	0126		C	A 19	94-2	1672	75	1994	0715	
AU 94	1736	503		A.	1	1995	0213		A	J 19:	94-7	3603		1994	0715	
AU 68	3514	18		В:	2	1998	0115									
EP 71	1029	93		A:	1	1996	0508		E	P 19	94-9	2253	3	1994	0715	

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R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL,
             PT, SE
                                        US 1993-92518
                                                            19930716
PRIORITY APPLN. INFO.:
                                        WO 1994-US7806
                                                            19940715
     A dry reagent prepd. by lyophilizing a labeled
AB
     ligand-immobilized receptor complex or a labeled
     receptor-immobilized ligand complex is, on rehydration, useful for
     detecting analytes in samples in a variety of displacement assays.
     Prepn. of a lyophilized support and use of
     lyophilized beads in a flow immunosensor are described, as
     is a lyophilization ELISA plate assay.
     50-99-7, Glucose, analysis
ΙT
     RL: ANT (Analyte); ANST (Analytical study)
        (lyophilized ligand-receptor complexes for assays and
        sensors)
L11 ANSWER 26 OF 37 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER:
                        1994:638436 CAPLUS
DOCUMENT NUMBER:
                         121:238436
                         Synthetic particulate vectors comprising a
TITLE:
                         non-liquid hydrophilic nucleus and amphiphilic
                         outer layer
                         Samain, Daniel; Delrieu, Pascal; Gibilaro,
INVENTOR(S):
                         Joelle; Dirson, Roselyne; Cervilla, Monique; De
                         Miguel, Ignacio; Ding, Li; Nguyen, Frederique;
                         Soulet, Nadine; Soler, Corinne
                         A et S Biovecteurs, Fr.
PATENT ASSIGNEE(S):
                         PCT Int. Appl., 45 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
                         French
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                      KIND
                            DATE
                                           APPLICATION NO.
                                                            DATE
                      ____
                                          _____
     ______
                            -----
                     A1
                            19940915
                                          WO 1994-FR228
                                                            19940301
     WO 9420078
         W: CA, JP, KR, US
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT,
                      A1
                            19940909
                                           FR 1993-2397
     FR 2702160
                                                            19930302
                            19950602
     FR 2702160
                      В1
                      AΑ
                            19940915
                                           CA 1994-2157384 19940301
     CA 2157384
                      A1
                            19951220
                                           EP 1994-908391
                                                            19940301
     EP 687173
                      В1
                            19970917
     EP 687173
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL,
             PT, SE
     JP 08507765
                       Т2
                            19960820
                                           JP 1994-519654
                                                            19940301
                                                          19940301
                      A1
                            19970709
                                           EP 1997-102586
     EP 782851
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL,
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                                                           19940301
                            19970806
                                           EP 1997-104152
     EP 787479
                       A1
           AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL,
             PT, SE
     AT 158179
                            19971015
                                           AT 1994-908391
                                                            19940301
                       E
     ES 2108432
                       Т3
                            19971216
                                           ES 1994-908391
                                                            19940301
     US 6013284
                            20000111
                                           US 1996-513853
                                                            19960501
                       Α
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Searcher: Shears 308-4994

FR 1993-2397

19930302

PRIORITY APPLN. INFO.:

EP 1994-908391 19940301 WO 1994-FR228 19940301

A synthetic particulate vector for pharmaceutical, cosmetic, or food AB prepns. comprises a non-liq. hydrophilic nucleus (e.g. a polysaccharide) and an outer layer at least partially consisting of amphiphilic compds. (e.g. phospholipids) and combined with the nucleus by hydrophobic interactions and/or ionic bonds. Amylopectin was mixed with 2N NaOH followed by addn. of a soln. of glycidyl trimethylammonium chloride in water and epichlorhydrine; the mixt. was then homogenized and the pH was set to 6 to obtain a gel which was washed and lyophilized. Thus, 0.6 g glucose oxidase (I) was mixed with 0.3 g above gel; the mixt. was then hydrated with 1.5 mL of buffer, pH = 7, and stirred at 4.degree. overnight and lyophilized. The above lyophilyzate was mixed with 0.15 g of hydrogenated soy phosphatidylcholine and 150 mL water, then it was homogenized to obtain I microcapsules with 92% microencapsulation.

L11 ANSWER 27 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1994:517757 CAPLUS

DOCUMENT NUMBER: 121:117757

TITLE: Synthesis of polymer bioactive conjugates

INVENTOR(S): Marcucci, Fabrizio; Gregory, Ruth
PATENT ASSIGNEE(S): Farmitalia Carlo Erba S.R.L., Italy

SOURCE: PCT Int. Appl., 44 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.					KIND DATE				1	APPLI	ο.	DATE				
	WO	9413	322			 1	1994		Ī				9	1993	1206		
		W:	AT,	AU,	BB,	BG,	BR,	BY,	CA,	CH,	CZ,	DE,	DK,	ES,	FI,	GB,	HU,
															PL,		
			-				UA,										
		RW:								GB.	GR,	IE,	IT,	LU,	MC,	NL,	PT,
															SN,		
	CA	21509															
	AU 9456968 AU 678796				В	2	1997	0612			•						
		6757]	EP 19	94-9	0269	2	1993	1206	
		6757															
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB.	GR,	ΙE,	IT,	LI,	LU,	MC,	NL,
			•	SE	·	·	•	•									
	JP	0850	4202		T	2	1996	0507			JP 19	93-5	1376	3	1993	1206	
		1682					1998	0815		1	AT 19	94-9	0269	2	1993	1206	
		2121					1998	1116]	ES 19	94-9	0269	2	1993	1206	
	US	6172	202		В	1	2001	0109		1	JS 19	97-8	8904	9	1997	0707	
PRIOR															1992	1204	
	CIONIII AILLICE INI									WO	1993-	EP34	29	W	1993	1206	
7.0	-		-														

AB A process for the prepn. of a conjugate between a polymer and a first substance having a biol. activity mediated by a domain consists of (a) contacting the first substance with a second substance which specifically binds to the domain of the first substance, (b) conjugating a polymer to the first substance having the second substance bound and (c) freeing the second substance from

the first substance having the polymer conjugate. The advantages such as prolonged half-life in vivo and reduced immunogenicity in proteins, that can be derived from the conjugation of polymers to drugs or diagnostic reagents are maintained. Thus, a PEG_monoclonal antibody conjugates (mAb 78) lyophilized formulation was prepd. contg. drug 0.05-0.5, excipient such as lactose or mannitol 2.5-5.0, surfactant (e.g., Poloxamer) 0.0025-0.025% (wt./vol.) and 6.5-7 pH-adjusting agent. The conjugate displayed a better retention of biol. activity than the unprotected conjugates.

L11 ANSWER 28 OF 37 CAPLUS COPYRIGHT 2001 ACS

1994:491424 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 121:91424

Removal/neutralization of hepatitis A virus TITLE:

during manufacture of high purity, solvent/

detergent factor VIII concentrate

Lemon, Stanley M.; Murphy, Paula C.; Smith, AUTHOR(S):

Andrew; Zou, Jinsheng; Hammon, John; Robinson,

Stephen; Horowitz, Bernard

Dep. Med., Univ. North Carolina, Chapel Hill, CORPORATE SOURCE:

NC, USA

J. Med. Virol. (1994), 43(1), 44-9 SOURCE:

CODEN: JMVIDB; ISSN: 0146-6615

DOCUMENT TYPE: Journal LANGUAGE: English

Recent reports have suggested an increased risk of type A viral hepatitis in hemophilic patients treated with high purity factor VIII concs. prepd. using ion exchange chromatog. coupled with solvent/detergent treatment for inactivation of viruses. To det. the capacity for removal or inactivation of hepatitis A virus during the factor VIII manufg. process, human plasma and various factor VIII prodn. intermediates were spiked with cell culture-propagated virus and subjected to scaled down conditions mimicking the manuf. of solvent/detergent factor VIII. The combination of antibody-mediated neutralization, cryopptn., anion exchange chromatog., and lyophilization in the absence of sucrose resulted in a minimal redn. of 5.5 to 8.55 log10 in the infectivity of hepatitis A virus.

L11 ANSWER 29 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1991:160306 CAPLUS

DOCUMENT NUMBER: 114:160306

TITLE: Immunoassay lyophilized reactant

mixture

INVENTOR(S): Cole, Francis X.

PATENT ASSIGNEE(S): Hygeia Sciences, Inc., USA

SOURCE: PCT Int. Appl., 32 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE ______ _____ -----WO 9013637 19901115 WO 1990-US2064 19900416 A1

W: CA, JP

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RW: AT, BE, CH, DE, DK, ES, FR, GB, IT, LU, NL, SE
                                                            19890428
    US 5102788
                      Α
                            19920407
                                           US 1989-344575
    CA 2053885
                      AA
                            19901029
                                           CA 1990-2053885
                                                            19900416
                                           EP 1990-908003
    EP 470192
                      Α1
                            19920212
                                                            19900416
    EP 470192
                      В1
                            19971008
        R: AT, BE, CH, DE, DK, ES, FR, GB, IT, LI, LU, NL, SE
                                           JP 1990-506814
     JP 05500854
                      T2
                            19930218
                                                            19900416
    AT 159047
                       Ε
                            19971015
                                           AT 1990-908003
                                                            19900416
    ES 2110415
                       Т3
                            19980216
                                           ES 1990-908003
                                                            19900416
PRIORITY APPLN. INFO.:
                                        US 1989-344575
                                                            19890428
                                        US 1985-747605
                                                            19850624
                                        US 1988-275656
                                                            19881121
                                        WO 1990-US2064
                                                            19900416
    A lyophilized mixt. of reactants for an immunoassay
AB
     includes antibody-gold sol particle conjugates,
    antibody latex particle conjugates, polyethylene glycol, a
    polyethylene glycol p-isooctylphenyl ether detergent and
     dextrin or trehalose. The polyethylene glycol is present
     to enhance binding of the immunoreactants and the polyethylene
     glycol p-isooctylphenyl ether detergent is present to
    prevent nonspecific interactions. The dextrin or threhalose
    prevents agglomeration of the polyethylene glycol and polyethylene
    glycol p-isooctylphenyl ether in the lyophilized mixt. at
    room temp. and facilitates retention of homogenous distribution of
    the ingredients of the mixt. to thereby enhance shelf life and
    redistribution of the mixt. in an aq. test system. Thus, an aq.
    mixt. to be subjected to lyophilization contained
    anti-human chorionic gonadotropin antibody conjugated to
    Au sol particles, a 2nd anti-human chorionic gonadotropin
    antibody conjugated to carboxylated modified latex
    particles, polyethylene glycol, Thimoseral, Tris buffer, Maltrin,
    disodium EDTA, and IGEPAl CA720.
ΙT
     99-20-7, Trehalose
    RL: ANST (Analytical study)
        (in immunoassay lyophilized reagent mixt. as
       antiagglomerant)
L11 ANSWER 30 OF 37
                     CAPLUS COPYRIGHT 2001 ACS
                         1990:527184 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         113:127184
                         Enzyme immunoassays and immunologic reagents for
TITLE:
                         home diagnostic application
                         Block, Elliott; Bahar, Izak; Cole, Frank; Eaton,
INVENTOR(S):
                         Cheryl A.; Jones, Wendy; Sigillo, Eric; Coseo,
                         Mary; Cicia, Nancy J.; Cannon, L. Edward;
                         Cantarow, Walter
PATENT ASSIGNEE(S):
                         Hygeia Sciences, Inc., USA
                         U.S., 15 pp. Cont. of U.S. Ser. No. 747,605,
SOURCE:
                         abandoned.
                         CODEN: USXXAM
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                           APPLICATION NO.
     PATENT NO.
                      KIND
                            DATE
                                                            DATE
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Searcher: Shears 308-4994

US 1988-275656

19881121

US 4931385

Α

19900605

US 5102788 19920407 US 1989-344575 19890428 Α US 1983-473907 19830310 PRIORITY APPLN. INFO.: US 1985-747605 19850624 US 1988-275656 19881121 Enzyme immunoassays, esp. ELISAs, for home diagnostic application AΒ under ambient room temp. and humidity, use a lyophilized mixt. contg. peroxidase-antibody conjugate, a binding-enhancer (e.g. PEG, polyvinyl alc., polyvinyl pyrrolidone, and dextran), a water-sol. nonionic surfactant in an amt. sufficient to provide detergency without having a deleterious effect on the conjugate, and a sugar (dextrin or trehalose). For an ELISA, a solid support is precoated with another antibody and then is treated with a blocking soln. comprising a blocking agent (bovine serum albumin, gelatin, milk proteins, or nonspecific IgG) and a water-sol. sugar. Both the lyophilized antibody conjugate mixt. and the immobilized antibody have preserved reactivity and immunolysis binding specificity even if exposed to high humidity and temps. of 80-120.degree.F prior to their use in the immunoassay. A diagnostic kit for the ELISA is disclosed. An ELISA for human chorionic gonadotropin (hCG) in urine used (1) lyophilized mixt. contg. peroxidase conjugated with a monoclonal antibody to the .beta.-chain of hCG, PEG, Hepes salt, Hepes acid, di-Na EDTA, MgSO4, dextrin, and IGEPAL CA-630 (octylphenoxypoly(ethyleneoxy)ethanol); (2) dipsticks coated with monoclonal antibody to hCG and treated with bovine serum albumin and sucrose in the blocking soln.; and (3) a chromogen soln. contg. tetramethylbenzidine, buffer, and H2O2. Urine was added to the conjugate mixt. and the dipstick was immersed in the soln. for >15 min. The dipstick was removed, washed with tap water, and dipped in the chromogen soln. for >5 min. When hCG was present, the dipstick changed from colorless to blue-green. 57-50-1, Sucrose, biological studies ITRL: BIOL (Biological study) (blocking soln. contg. bovine serum albumin and, for stable antibody-coated dipstick for home diagnostic ELISA) IT 99-20-7, Trehalose RL: BIOL (Biological study) (heat- and humidity-stable lyophilized mixt. contg. peroxidase-antibody conjugate and, for home diagnostic ELISA) L11 ANSWER 31 OF 37 CAPLUS COPYRIGHT 2001 ACS 1989:436239 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 111:36239 Device and method for biochemical assay TITLE: INVENTOR(S): Stanley, Christopher John; Johannsson, Axel PATENT ASSIGNEE(S): IQ (BIO) Ltd., UK SOURCE: PCT Int. Appl., 49 pp. CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: KIND DATE APPLICATION NO. DATE PATENT NO. ----------------- _____ WO 8804428 **A**1 19880616 WO 1987-GB899 19871211

W: AU, DK, JP, US

RW: AT, BE, CH, DE, FR, GB, IT, LU, NL, SE

AU 8783339 A1 19880630 AU 1987-83339 19871211
DK 8804488 A 19880811 DK 1988-4488 19880811
PRIORITY APPLN. INFO.: GB 1986-29740 19861212
WO 1987-GB899 19871211

An app. for carrying out a biochem. assay, e.g. an immunoassay or AB nucleic acid hybridization assay, comprises a hard polystyrene reaction surface adapted to bind a 1st biochem. ligand (e.g. a monoclonal antibody), a liq. absorbent wadding adjacent to the reaction surface to absorb washing soln. applied to the reaction surface, a filter matrix overlying the reaction surface in close contact therewith for filtering a sample and for retaining a 2nd biochem. ligand in contact with the reaction surface, wherein the 2nd ligand can specifically bind to the 1st ligand on the surface. The filter matrix is removable to facilitate washing of the reaction surface, and the filter matrix comprises a labeled substance capable of a specific binding reaction at the reaction surface during the assay. Polystyrene strips (for reaction surfaces) were dipped into monoclonal anti-Chlamydia antibody (5 .mu.g/mL) and incubated overnight at 37.degree. before dipping in glazing soln. (lactose, degraded gelatin, thimerosal, and Tween 20) and air drying. Latex-filled paper (filter matrix) was preblocked by immersion for 1 h in the glazing soln., and a soln. contg. Fab' fragments of anti-Chlamydia antibody conjugated to alk. phosphatase was pipetted onto 10-mm2 paper squares. The conjugate was freeze-dried into the filter matrix. A liq. sample contg. Chlamydia antigen was applied to the filter matrix on the polystyrene reaction surface, the reaction surface was washed with buffer contg. diethanolamine, iodonitrotetrazoleum violet, NaN3, EDTA, and EtOH, and developed using a developer pad contg. Cu-activated pig heart diaphorase, alc. dehydrogenase, sucrose, Triton X-705, gelatin, and Tris. A red color appeared on the developer pad within 5 min.

L11 ANSWER 32 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1989:73606 CAPLUS

DOCUMENT NUMBER: 110:73606

TITLE: Cytotoxic liposomes: membrane interleukin 1

presented in multilamellar vesicles

AUTHOR(S): Bakouche, Ouahid; Lachman, Lawrence B.; Knowles,

Rebecca D.; Kleinerman, Eugenie S.

CORPORATE SOURCE: Dep. Cell Biol., M. D. Anderson Hosp., Houston,

TX, 77030, USA

SOURCE: Lymphokine Res. (1988), 7(4), 445-56

CODEN: LYREDH; ISSN: 0277-6766

DOCUMENT TYPE: Journal LANGUAGE: English

AB Paraformaldehyde-fixed lipopolysaccharide (LPS)-activated human monocytes produced significant lysis of the human melanoma cell line A375. The cytotoxic activity was retained following treatment of the fixed monocytes with anti-tumor necrosis factor (anti-TNF) antibodies but was specifically inhibited by a mixt. of anti-TNF and anti-interleukin 1 (anti-IL-1) antibodies. A375 cells were also killed by plasma membranes purified from LPS-activated human blood monocytes. This activity was specifically inhibited by anti-IL-1 .alpha. antibodies, but only partially inhibited by anti-IL-1 .beta.

antibodies. CHAPS detergent-extd. plasma-membrane IL-1 in its sol. form or assocd. with lyophilized liposomes was also able to kill A375 cells, and this antitumor activity was inhibited by anti-IL-1 antibodies. These results suggest that membrane IL-1, primarily IL-1 .alpha., was cytotoxic for the A375 cells. CKS-17, a peptide synthesized with homol. to a highly conserved region of the immunosuppressive retroviral envelope protein P15E, when covalently bound to BSA, partially inhibited the IL 1 activities of tumor cell cytotoxicity and T-cell clone proliferation, displayed by purified plasma membranes, detergent-extd. membrane IL 1, or membrane IL 1 assocd. with liposomes. Thus, cytotoxic membrane IL-1 can be solubilized by detergent, bound to the surface of liposomes, and specifically inhibited by anti-IL-1 antibodies or the immunosuppressive peptide CKS-17.

L11 ANSWER 33 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1988:545719 CAPLUS

DOCUMENT NUMBER: 109:145719

TITLE: Enzyme immunoassay-monoclonal test system for

the detection of plague pathogens

AUTHOR(S): Temiralieva, G. A.; Arakelyan, I. S.; Lukhnova,

L. Yu.; Apsatarova, R. A.

CORPORATE SOURCE: Sredneaziat. Nauchno-Issled. Protivochumn.

Inst., Alma-Ata, USSR

SOURCE: Lab. Delo (1988), (8), 61-3

CODEN: LABDAZ; ISSN: 0023-6748

DOCUMENT TYPE: Journal LANGUAGE: Russian

AB EIA based on the use of monoclonal antibodies (isolated from ascitic fluid), described for detecting plague pathogen under lab. and field conditions, uses antibody-horseradish peroxidase conjugates and Tween as the reaction stabilizer. The conjugates were prepd. by periodate method. Caprylic acid gave the most pure plague monoclonal Igs, the diln. of the conjugates was .gtoreq.1:1000. One part Ig and 2 parts of the enzyme was the best ratio. This conjugate ratio can detect 30 ng/mL plague microbes. Medium contg. polyvinylpyrrolidone with 7.5% sucrose was the best for lyophilizing the conjugates. The lyophilized conjugates preserved their sp. activity for 2 yr at -20.degree. to 25.degree.. The EIA had

L11 ANSWER 34 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1986:107939 CAPLUS

DOCUMENT NUMBER: 104:107939

TITLE: Preparation of pure yeast chromatin

higher sensitivity than the agglutination test.

INVENTOR(S): Shalitin, Channa

PATENT ASSIGNEE(S): Technion Research and Development Foundation

Ltd., Israel

SOURCE: Ger. Offen., 27 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent
LANGUAGE: German

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

DE 1985-3525722 19850718

19840719

IL 1984-72452

19860123

A1 19880429

A1

DE 3525722

IL 72452

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IL 1984-82930
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    GB 2163165
                     B2
                           19880720
    GB 2163165
                                      IL 1984-72452
                                                          19840719
PRIORITY APPLN. INFO.:
    Yeast chromatin is purified by d. gradient centrifugation of the
    cell homogenate. Thus, baker's yeast cells were disrupted with
    glass beads. Phenylmethylsulfonyl fluoride (I) was added and the
    mixt. centrifuged. The chromatin pellet was suspended in pH 7.5
    buffer contq. I, a surfactant, and an antioxidant and
    centrifuged. The pellet was suspended and centrifuged through a
    sucrose soln. Purified chromatin was treated with a
    nuclease in buffer contg. I and centrifuged. The supernatant was
    centrifuged on a 5-20% sucrose gradient and sep. fractions
    were dialyzed and lyophilized to obtain p20 protein. The
    p20 protein may be injected, with an adjuvant, into lab animals to
    elicit the formation of antibodies to ras oncogene
    proteins.
ΙT
    57-50-1, biological studies
    RL: BIOL (Biological study)
        (in chromatin purifn., from yeast)
L11 ANSWER 35 OF 37 CAPLUS COPYRIGHT 2001 ACS
                        1984:412196 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                        101:12196
                        Gamma globulin-containing compositions
TITLE:
INVENTOR(S):
                        Hooper, John A.; Mankarious, Samia; Liu-Rash,
                        Catherine R.
PATENT ASSIGNEE(S):
                        Baxter Travenol Laboratories, Inc., USA
                        PCT Int. Appl., 21 pp.
SOURCE:
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                        English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
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    WO 8400891
                    A1
                           19840315
                                         WO 1983-US1016
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PRIORITY APPLN. INFO.:
                                       US 1982-413059
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                                       EP 1983-902407
                                                          19830701
                                       WO 1983-US1016
AB
    The anticomplement activity of .gamma.-globulin products prepd. by
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Searcher :

Shears 308-4994

methods involving ultrafiltration and(or) treatment with ion exchangers is considerably decreased by including .gtoreq.1 nonsurface-active stabilizers such as hydrophilic macromols., amino acids, and low mol. wt. polyols with .gamma.-globulin concns. during ultrafiltration and(or) ion exchange. An aq. soln. of Cohn Fraction II at a protein concn. of 50 g/L and pH 5.3 was dialyzed against 4 vols. of a soln. contg. 20 mM NaCl and 0.5 g/L PEG-4000 [25322-68-3]. The Fraction II soln. was pumped through Millipore cassettes where materials with mol. wt. of <100,000 Daltons were removed by ultrafiltration. The retentate which contained concd. Ig was returned to and mixed with Ig soln. undergoing dialysis. The dialysis soln. was continuously pumped into the Iq soln. at the same rate as filtrate was produced and removed. When 300 L dialysis soln. was depleted, the Ig soln. was concd. to a protein concn. of 55 mg/mL, and the concd. Ig soln. treated with hydrated DEAE Sephadex A-50 (5 g/g protein), agitated for 3 h at 5.degree., and filtered. NaCl, dextrose [50-99-7], glycine [56-40-6], and albumin were added to the soln. to a final concn. of 0.85, 2.0, 2.25, and 0.1%, resp., the pH adjusted to 7.0 and protein concn. adjusted to 5.2%. The soln. contained .apprx.1.3 mg PEG-4000/mL from the dialysis step. The soln. was sterile filtered, filled into vials, lyophilized and the vial sealed.

50-99-7, biological studies ΙT RL: BIOL (Biological study)

(anticomplement activity of .gamma.-globulin concs. decreased by)

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Possibility of producing antigens from TITLE: Histoplasma capsulatum using detergents

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Antigens were extd. from freeze-dried mycelium of H. capsulatum with cetyltrimethylammonium bromide (I), Na glycocholate (II), Na oleate (III), sulfanol (IV), and Tween 80 (V). Obtained antigenic fractions were mainly polysaccharide and had a low nucleic acid content (0.1-0.4%). The protein content depended on the type of detergent used in the extn. procedure. Antigens extd. with I, II, III, IV, and V contained 17.5, 20, 36.2, 42.5, and 24% protein resp. For immunol. characterization the antigenic activity of fractions was compared with that of the antigen obtained by the conventional .beta.-naphthol method. According to passive hemagglutination and antibody neutralization reactions, antigen fractions obtained by extn. with IV and I were at least as active as that prepd. by the .beta.-naphthol method. A lower antigen content was found in fractions obtained by extn. with II, III, and esp. with V. All fractions showed 2 pptg. zones in gel with serums of hyperimmunized rabbits. Antigen extn. with the